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(54) The gal operon of streptomyces.

(57) A recombinant DNA molecule comprising the Streptomyces gal operon galK gene; galE gene; galT gene; P1 promoter; P2 promoter; P2 promoter expression unit; P1 promoter regulated region; or the entire Streptomyces gal operon is prepared.

TITLE THE GAL OPERON OF

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STREPTOMYCES

CROSS REFERENCE TO RELATED APPLICATIONS

This application is a continuation-in-part of Serial Number 834,706, filed February 28, 1986, which is pending.

BACKGROUND OF THE INVENTION

This invention relates to a recombinant DNA molecule comprising the <u>Streptomyces gal</u> operon.

Hodgson, J. Gen. Micro., 128, 2417-2430 (1982), report that Streptomyces coelicolor A3(2) has a glucose repression system which allows repression at the level of transcription of the arabinose uptake system, one of the glycerol uptake systems, and also repression of the galactose uptake system in wild type strains. There is no report in Hodgson of actual galactose metabolism by \underline{S} . coelicolor A3(2).

Okeda et al. Mol. Gen. Genet., 196, 501-507

(1984), report that glucose kinase activity, 2-deoxyglucose-sensitivity, glucose utilization and glucose repression were all restored to S. coelicolor A3(2) glk (glucose kinase) mutants transformed by a 3.5 kb DNA

fragment which contained the glk gene cloned from S.

coelicolor into a phage vector.

Seno et al., Mol. Gen. Genet., 193, 119-128 (1984), report the glycerol (gyl) operon of Streptomyces coelicolor, and state that such operon is substrate-inducible and catabolite-repressible.

Debouck et al., Nuc. Acids. Res., 13(6), 1841-1853 (1985), report that the gal operon of E. coli consists of three structurally contiguous genes which specify the enzymes required for the metabolism of galactose, i.e., galE (uridine diphosphogalactose-4-epimerase), galT (galactose-1-phosphate uridyltransferase) and galK (galactokinase); that such genes are expressed from a polycistronic mRNA in the order E, T, K; that the expression of the promoter distal gene of the operon, galK, is known to be coupled translationally to the galT gene immediately preceding it; that such translational coupling results from a structural overlap between the end of the galT coding sequence and the ribosome binding region of galK; and that the translational coupling of galT and galK ensures the coordinate expression of these genes during the metabolism of galactose.

SUMMARY OF THE INVENTION

This invention relates to a recombinant DNA molecule comprising a <u>Streptomyces gal</u> operon <u>gal</u>K gene; <u>gal</u>E gene; <u>gal</u>T gene; P2 promoter expression unit, or P2 promoter or any functional derivative thereof as well as a recombinant DNA molecule comprising a <u>Streptomyces gal</u> operon P1 promoter, P1 promoter regulated region or the entire <u>gal</u> operon or any regulatable and functional derivative thereof.

This invention also relates to a recombinant DNA molecule comprising the <u>Streptomyces gal</u> operon or any regulatable and functional derivative thereof and a functional DNA molecule operatively linked to such operon; a recombinant DNA vector comprising and such DNA molecule, and, optionally, additionally comprising a replicon; a method of preparing a host cell transformed with such

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vector; the transformed host prepared by such method; a method of expressing such functional DNA sequence which comprises cultivating such transformed host under suitable conditions such that the functional DNA sequence is expressed; and to a method of regulating the expression of such functional DNA sequence which comprises cultivating such transformed host under conditions which regulate such expression.

This invention also relates to a recombinant DNA molecule comprising the <u>Streptomyces gal</u> operon P2 promoter expression unit or any functional derivative thereof and a functional DNA molecule operatively linked to such unit; a recombinant DNA vector comprising such DNA molecule, and, optionally, additionally comprising a replicon; a method of preparing a host cell transformed with such vector; the transformed host prepared by such method; and to a method of expressing such functional DNA sequence which comprises cultivating such transformed host under suitable conditions such that the functional DNA sequence is expressed.

This invention also relates to a recombinant DNA molecule comprising the <u>Streptomyces gal</u> operon Pl promoter regulated region or any regulatable and functional derivative thereof and a functional DNA molecule operatively linked to such region; a recombinant DNA vector comprising such DNA molecule, and, optionally, additionally comprising a replicon; a method of preparing a host cell transformed with such vector; the transformed host prepared by such method; a method of expressing such functional DNA sequence which comprises cultivating such transformed host under suitable conditions such that the functional DNA sequence is expressed; and to a method of regulating the expression of such functional DNA sequence which comprises cultivating such transformed host under conditions which regulate such expression.

This invention also relates to a recombinant DNA

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molecule comprising the Streptomyces gal operon Pl 1 promoter or any regulatable and functional derivative thereof and a foreign functional DNA molecule operatively linked to such region; a recombinant DNA vector comprising such DNA molecule, and, optionally, additionally 5 comprising a replicon; a method of preparing a host cell transformed with such vector; the transformed host prepared by such method; a method of expressing such functional DNA sequence which comprises cultivating such transformed host under suitable conditions such that the 10 functional DNA sequence is expressed; and to a method of regulating the expression of such functional DNA sequence which comprises cultivating such transformed host under conditions which regulate such expression.

This invention also relates to a recombinant DNA molecule comprising the Streptomyces gal operon P2 promoter or any functional derivative thereof and a foreign functional DNA molecule operatively linked to such region; a recombinant DNA vector comprising such DNA molecule, and, optionally, additionally comprising a replicon; a method of preparing a host cell transformed with such vector; the transformed host prepared by such method; and to a method of expressing such functional DNA sequence which comprises cultivating such transformed host under suitable conditions such that the functional DNA sequence is expressed.

This invention also relates to a method of enabling a non-galactose utilizing host microorganism or cell to utilize galactose which comprises transforming such host with a recombinant DNA molecule comprising a Streptomyces gal operon or any portion of the Streptomyces gal operon, or any functional derivative thereof, which is adequate to enable such transformed host to utilize galactose. This invention also relates to the recombinant DNA vector employed in such method and to the host prepared by such method.

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BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 represents a restriction endonuclease map of the <u>Streptomyces lividans</u> 1326 galactose (gal) operon and indicates approximate locations for structural genes and promoters within the operon.

Figure 2 represents a restriction endonuclease map of plasmid pK21.

Figure 3 represents a comparison of the restriction endonuclease maps of the <u>S. lividans gal</u> operon and a restriction fragment containing the <u>S</u>. coelicolor galk gene.

DETAILED DESCRIPTION OF THE INVENTION

genome contains a operon for the metabolism of galactose
(i.e., a gal operon) which comprises three structural
genes (galT, galE and galK) and two promoters (Pl and
P2). The galT gene product is known as galactose-lphosphate uridyltransferase (transferase), the galE gene
product is known as uridine diphosphogalactose4-epimerase (epimerase), and the galK gene product is
known as galactose-l-kinase (galactokinase). The function
of the gene products of galT, galE and galK in galactose
metabolism in Streptomyces is explained by the following
diagram:

- 1. galactose + ATP galactokinase
 galactose-1-phosphate + ADP
- galactose-l-phosphate + UDP-glucose transferase
 UDP-galactose + glucose-l-phosphate
- 3. UDP-qalactose epimerase UDP-glucose

By the term "promoter" is meant any region upstream of a structural gene which permits binding of RNA polymerase and transcription to occur.

By the term "structural gene" is meant a coding 1 sequence for a polypeptide which serves to be the template for the synthesis of mRNA.

By the term "operon" is meant a group of closely linked genes responsible for the synthesis of one or a group of enzymes which are functionally related as members of one enzyme system. An operon comprises an operator gene, a number of structural genes (equivalent to the number of enzymes in the system) and a regulator gene. Ву "operator" or "operator gene" is meant a DNA sequence which controls the biosynthesis of the contiguous structural gene(s) within an operon. By "regulator gene" is meant a gene which controls the operator gene in an operon through the production of a repressor which can be either active (enzyme induction) or inactive (enzyme repression). The transcription of the structural gene(s) in an operon is switched on or off by the operator gene which is itself controlled in one or more of three ways: 1) in inducible enzyme systems, the operator is switched off by a repressor produced by the regulator gene and 20 which can be inactivated by some metabolite or signal substance (an inducer) coming from elsewhere in the cell or outside the cell, so that the presence of the inducer results in the operon becoming active; or 2) in repressed enzyme systems, the operator is switched off by a 25 repressor-corepressor complex which is a combination of an inactive repressor produced by the regulator gene with a corepressor from elsewhere, so that the presence of the corepressor renders the operon inactive; or 3) in activated gene systems, the promoter is switched on by an 30 activator produced by a regulator gene which can be activated by some metabolic or signal substance.

The Streptomyces gal operon is naturally present in the Streptomyces genome.

By the term "Streptomyces gal operon" is meant that region of the Streptomyces genome which comprises the

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Pl promoter, P2 promoter, <u>qal</u>T, <u>qal</u>E and <u>qal</u>K structural genes and any other regulatory regions required for transcription and translation of such structural genes.

By the term "regulatory region" is meant a DNA sequence, such as a promoter or operator, which regulates transcription of a structural gene.

The following model is suggested for gene expression within the <u>Streptomyces gal</u> operon. The Pl promoter is a galactose inducible promoter (i.e., it is induced in the presence of galactose and repressed in the presence of glucose). According to Sl data, the P2 promoter is constitutive, i.e., it is "turned on" regardless of the presence or absence of galactose or any other carbon source.

15 A cosmid library was constructed for Streptomyces lividans 1326 DNA by using cosmid pJW357 (which encodes the ability to replicate in both Streptomyces and E. coli). This library was then transfected into E. coli K21 which is a derivative of the E. coli strain MM294 which 20 contained a bacteriophage Pl transduced galactokinase (galK) mutation. Transfected cells were plated under media conditions which select for both the presence of the cosmid and the presence of an active galk gene. Weakly positive colonies were isolated and the cosmid DNA derived 25 from these colonies was transformed into the K21 strain. These transformations yielded two cosmids which consistently produced positive growth with galactose as the only carbon source. These galK cosmids were then transformed into a Streptomyces host (i.e., Streptomyces 30 lividans 1326-12K) which had been isolated by the inventors of the subject invention as unable to grow on medium in which galactose was the only carbon source by using 2-deoxy-galactose selection [see, Brawner et al., Gene, 40 191 (1985), in press]. Under conditions which differentiate strains able and unable to produce 35

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galactokinase, only one of the cosmids caused the

Streptomyces lividans 1326-12K host to become galk⁺.

Further studies have demonstrated that this cosmid encodes
a gene with galactokinase activity. Additional studies,
including DNA sequence analysis and protein studies
demonstrate that this Streptomyces gene shares homology
with the E. coli and yeast galactokinase genes.
Regulation studies indicate that the cosmid encoded
galactokinase gene regulated in the same manner as the
chromosome encoded gene.

A. S. lividans gal operon was originally isolated from a ca. 9 kilobase (Kb) region of Streptomyces lividans 1326. The ca. 9 Kb region of Streptomyces lividans 1326 containing the Streptomyces gal operon has been mapped 15 substantially as follows in Table A. By "substantially" is meant (i) that the relative positions of the restriction sites are approximate, (ii) that one or more restriction sites can be lost or gained by mutations not otherwise significantly affecting the operon, and (iii) 20 that additional sites for the indicated enzymes and, especially for enzymes not tested, may exist. restriction enzymes used herein are commercially available. All are described by Roberts, Nuc. Acids. Res., 10(5): p117 (1982). 25

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TABLE A

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	Map Position	Restriction Enzyme	Location (kb)
5	1	<u>Hind</u> III	40
	1a	<u>Nru</u> I	. 0
	2	<u>Bgl</u> II	.75
	3	ECORI	1.05
	4	<u>Pvu</u> II	1.15
10	5	<u>Mlu</u> I	2.30
	6	<u>Pvu</u> II	2.80
	7	ECORI	4.00
	8	PvuII	4.10
	8 a	<u>Sac</u> I	4.25
15	9	PvuII	5.00
	10	<u>Xho</u> I	5.50
	11	<u>BamH</u> I	5.80
	12	<u>BamH</u> I	6.50
	13	MluI	6.90
20	13a	PvuII	7.20
	14	MluI	7.80
	15	<u>BamH</u> I	8.00
	16	<u>Sph</u> I	8.30

25 Figure 1 represents a restriction endonuclease map of the <u>Streptomyces lividans</u> 1326 <u>gal</u> operon and indicates locations for structural genes (<u>gal</u>T, <u>gal</u>E and <u>gal</u>K) and promoters (P1 and P2) comprised within the operon.

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Referring to Table A and Figure 1, the location of the promoters and structural genes of the <u>Streptomyces</u>

<u>lividans</u> 1326 <u>gal</u> operon are mapped substantially as follows in Table B:

TABLE B

		Location (Kb)
	Pl transcription start site	.10
10	galT translation initiation codon	.15
	P2 transcription start site	1.25
	galE translation initiation codon	1.50
	galK translation initiation codon	2.40
	3' end of galk message	3.60
TD		

Microorganisms of the genus <u>Streptomyces</u> have historically been used as a source of antibiotics for the pharmaceutical industry. Consequently, the technical skills necessary to scale-up the production of biological products using <u>Streptomyces</u> as the vehicle for the production of such products are presently available. However, before <u>Streptomyces</u> can be used as a vehicle for the production of bioactive molecules using the new recombinant DNA technologies, there is a need to define regulatory elements in <u>Streptomyces</u> analogous to those which have proved useful in <u>E. coli</u>. These regulatory elements include ribosomal binding sites and regulated transcriptional elements.

The existence of a galE, galT or galK gene or gene product or gal operon in Streptomyces has not been previously reported. The instant invention, i.e., the cloning of the Streptomyces gal operon, enables construction of regulatable expression/cloning vectors in Streptomyces, other actinomycetes, and other host organisms. Furthermore, the instant invention led to the discovery that the Streptomyces gal operon is

- 1 polycistronic. Perhaps the most important feature of the cloning of the <u>Streptomyces gal</u> operon is the observation that there are sequences essential for regulation of the <u>Streptomyces gal</u>K gene. Direct analogy to the initial use
- of the <u>lac</u> promoter from <u>E. coli</u> as an expression system can be made. In fact, Brosius et al., <u>Proc. Natl. Acad.</u>
 <u>Sci. USA</u>, <u>81</u>, 6929-6933 (1984), utilized the regulatory elements of the <u>E. coli</u> lac promoter to regulate the exceptionally strong <u>E. coli</u> ribosomal promoters. Because
- 10 it is likely that the <u>Streptomyces gal</u> operon ribosomal promoters are also exceptionally strong, such promoters enable the construction of regulatable expression vectors which will be very useful in <u>Streptomyces</u>, other actinomycetes, and other host organisms. The instant
- 15 invention also enabled the unexpected discovery that the 2-deoxygalactose selection which has been used in <u>E. coli</u> to select for <u>galk</u> mutants also operates in <u>Streptomyces</u> to select for <u>galk</u> mutants [see, Brawner et al., <u>Gene 40</u>, 191 (1985), in press]. This observation, combined with
- 20 the ability to clone the <u>Streptomyces galk</u> gene and the promoter and regulatory regions required for its transcription and translation on a cosmid, as described herein, allows the direct insertion of any structural gene into the chromosomally located galk gene of <u>Streptomyces</u>
- 25 by homologous recombination. This manipulation will allow molecular biologists to stably insert DNA fragments of interest into the Streptomyces chromosome. Such an approach will allow researchers to tag or mark a Streptomyces strain of interest or to insert expression
- 30 cassettes into the organism without the need of maintaining an antibiotic selection such as that presently required by most <u>Streptomyces</u> expression vectors.

This invention relates to a recombinant DNA molecule comprising the <u>Streptomyces gal</u> operon or any 35 regulatable and functional derivative thereof.

- l By "regulatable and functional derivative" is meant any derivative of the <u>Streptomyces gal</u> operon which functions in substantially the same way as the naturally occurring <u>Streptomyces gal</u> operon in terms of regulatable production
- of the galT, galE and galK gene products. Such derivatives include partial sequences of the gal operon, as well as derivatives produced by modification of the gal operon coding sequence. Techniques for modifying the gal operon which are known in the art include, for example,
- 10 treatment with chemical mutagens, irradiation or direct genetic engineering, such as by inserting, deleting or substituting nucleic acids by the use of enzymes or recombination techniques. The naturally occurring Streptomyces gal operon can be isolated from any galactose
- 15 utilizing Streptomyces strain by employing the techniques described herein. Numerous strains of various Streptomyces species are publicly available from many sources. For example, the American Type Culture Collection, Rockville, Maryland, U.S.A. has approximately
- 20 400 different species of Streptomyces available to the public. The ability of a particular strain of Streptomyces to utilize galactose can be readily determined by conventional techniques, such as by growing such strain on a medium containing galactose as the sole
- which to isolate a gal operon include S. lividans, S. coelicolor, S. azuraeus and S. albus, S. carzinostaticus, S. antifibrinolyticus and S. longisporus. S. lividans is most preferred. The Streptomyces gal operon, and smaller
- 30 portions thereof, is useful as a nucleic acid probe to obtain homologous sequences from other cells and organisms. The <u>Streptomyces gal</u> operon is also useful as a selection marker in an appropriate host mutant, and for providing regulatory elements. By "appropriate host
- 35 mutant " is meant a host which does not utilize galactose

- because it (a) does not contain a gal operon or (b) contains a nonfunctional gal operon, or (c) contains a defect within a homologous structural gene or regulatory region comprised by the <u>Streptomyces gal</u> operon such as a
- defective Pl promoter, P2 promoter, galT gene, galK gene and/or galE gene. Thus, a recombinant DNA molecule (comprising the Streptomyces gal operon and a foreign functional DNA sequence operatively linked thereto), which can be prepared by conventional techniques, can be
- 10 transformed into an appropriate host mutant by conventional techniques for incorporation into the host genome by homologous recombination to enable regulatable expression of the foreign functional DNA sequence without the need of maintaining an expensive antibiotic
- on recombinant DNA expression vectors for regulatable expression of a foreign functional DNA sequence operatively linked to such operon in an appropriate host mutant transformed with such vector without the need of
- 20 maintaining an expensive antibiotic selection. Such operon is also useful for transforming those cells, viruses and microorganisms, such as strains of Streptomyces, other actinomycetes, and other prokaryotic organisms, such as gal E. coli strains, which do not
- 25 utilize galactose into galactose utilizing strains. Such transformation may have pleiotrophic effects on the transformed host. By the term "functional DNA sequence" is meant any discrete region of DNA derived directly or indirectly from Streptomyces or any other source which
- gene expression unit, structural gene, promoter or a regulatory region. Preferred functional DNA sequences include those coding for polypeptides of pharmaceutical importance, such as, but not limited to, insulin, growth
- 35 hormone, tissue plasminogen activator, alpha -l-antitrypsin or antigens used in vaccine production. By the

term "foreign functional DNA sequence" is meant a functional DNA sequence not derived from the Streptomyces gal operon coding region.

This invention also relates to a recombinant DNA molecule comprising the Streptomyces gal operon P2 5 promoter expression unit or any functional derivative thereof. By the term "P2 promoter expression unit" is meant that region of the Streptomyces gal operon comprising the Streptomyces gal operon P2 promoter, galE and galk structural genes and any other regulatory regions required for transcription and translation of such structural genes. By "functional derivative" is meant any derivative of the Streptomyces gal operon P2 promoter expression unit which functions in substantially the same way as the naturally occurring region in terms of production of the Streptomyces gal operon galE and galK gene products. Such derivatives include partial sequences of the Streptomyces gal operon P2 promoter expression unit, as well as derivatives produced by modification of the Streptomyces gal operon P2 promoter expression unit 20 coding sequence. Techniques for effecting such modification are known in the art, and some have been outlined above. The naturally occurring Streptomyces gal operon P2 promoter expression unit can be isolated from the naturally occurring Streptomyces gal operon by 25 conventional techniques. The Streptomyces gal operon P2 expression unit is useful as a selection marker in an appropriate host mutant and for providing regulatory elements. By "appropriate host mutant" is meant a host which does not utilize galactose because it contains a defect within a homologous structural gene or regulatory region comprised by the Streptomyces P2 promoter expression unit such as a defective P2 promoter, galE gene and/or galk gene. Thus, a recombinant DNA molecule (comprising the Streptomyces gal operon P2 promoter expression unit and a foreign functional DNA sequence

- operatively linked thereto), which can be prepared by conventional techniques, can be transformed into an appropriate host mutant by conventional techniques for incorporation into the host genome by homologous
- recombination to enable constitutive expression of the foreign functional DNA sequence without the need of maintaining an expensive antibiotic selection. Such expression unit may also be incorporated on recombinant DNA expression vectors for constitutive expression of
- 10 foreign functional DNA sequences. The Stretomyces gal operon P2 promoter expression unit is also useful for complementation of an appropriate host mutant which can then be used for constitutive expression of a foreign functional DNA sequence operatively linked to such
- 15 expression unit in an appropriate host mutant transformed with such vector without the need of maintaining an expensive antibiotic selection.

This invention also relates to a recombinant DNA molecule comprising the Streptomyces gal operon Pl

20 promoter regulated region or any regulatable and functional derivative thereof. By the term "Pl promoter regulated region" is meant that region of the Streptomyces gal operon Pl promoter, galT, galE and galK structural genes and any

25 other regulatory regions required for transcription and translation of such structural genes. By "regulatable and functional derivative" is meant any derivative of the Streptomyces gal operon Pl promoter regulated region which functions in substantially the same way as the naturally occurring region in terms of regulatable production of the Streptomyces gal operon galT, galE and galK gene products. Such derivatives include partial sequences of

as well as derivatives produced by modification of the 35 <u>Streptomyces gal</u> operon Pl promoter regulated region coding sequence. Techniques for effecting such

the Streptomyces gal operon Pl promoter regulated region,

modification are known in the art, and some have been outlined above. The naturally occurring Streptomyces gal 1 operon Pl promoter regulated region can be isolated from the naturally occurring Streptomyces gal operon by conventional techniques, such as by excising the P2 promoter from the naturally occurring Streptomyces gal 5 operon or inactivating the P2 promoter by a point mutation or by inserting a foreign DNA sequence within the promoter. The Streptomyces gal operon Pl promoter regulated region is useful for the utilities outlined above for the Streptomyces gal operon. 10

This invention also relates to a recombinant DNA molecule comprising the Streptomyces gal operon P2 promoter or any functional derivative thereof. "functional derivative" is meant any derivative of the Streptomyces gal operon P2 promoter which functions in substantially the same way as the naturally occurring P2 promoter in terms of enabling the binding of RNA polymerase thereto and transcription of a functional DNA sequence operatively linked to such promoter. derivatives include partial sequences of the Streptomyces gal operon P2 promoter, as well as derivatives produced by modification of the gal operon P2 promoter coding sequence. Techniques for effecting such modification are known in the art, and some have been outlined above. naturally occurring Streptomyces gal operon P2 promoter 25 can be isolated from the naturally occurring Streptomyces gal operon by conventional techniques. A recombinant DNA molecule (comprising the Streptomyces gal operon P2 promoter and a foreign functional DNA sequence operatively linked thereto), which can be prepared by conventional 30 techniques, can be transformed into an appropriate host mutant by conventional techniques for incorporation into the host genome by homologous recombination to enable constitutive expression of the foreign functional DNA sequence. The Streptomyces gal operon P2 promoter is also 35

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- l useful for incorporation into recombinant DNA expression vectors for constitutive expression of a foreign functional DNA sequence operatively linked thereto in viruses and eukaryotic or prokaryotic cells or organisms,
- 5 especially in <u>Streptomyces</u> or other actinomycetes, transformed with such vector.

This invention also relates to a recombinant DNA molecule comprising the <u>Streptomyces gal</u> operon Pl promoter or any regulatable and functional derivative

- 10 thereof. By "regulatable and functional derivative" is meant any derivative of the <u>Streptomyces gal</u> operon Pl promoter which functions in substantially the same way as the naturally occurring Pl promoter in terms of enabling the binding of RNA polymerase thereto and regulating the
- 15 transcription of a functional DNA sequence operatively linked to such promoter. Such derivatives include partial sequences of the <u>Streptomyces gal</u> operon Pl promoter, as well as derivatives produced by modification of the <u>gal</u> operon Pl promoter coding sequence. Techniques for
- 20 effecting such modification are known in the art, and some have been outlined above. The naturally occurring Streptomyces gal operon Pl promoter can be isolated from the naturally occurring Streptomyces gal operon by conventional techniques. A recombinant DNA molecule
- 25 (comprising the <u>Streptomyces gal</u> operon Pl promoter and a foreign functional DNA sequence operatively linked thereto), which can be prepared by conventional techniques, can be transformed into an appropriate host mutant by conventional techniques for incorporation into
- the host genome by homologous recombination to enable regulatable expression of the foreign functional DNA sequence. The <u>Streptomyces gal</u> operon Pl promoter is also useful for incorporation into recombinant DNA expression vectors for regulatable expression of a foreign functional
- 35 DNA sequence operatively linked thereto in viruses and eukaryotic or prokaryotic cells or organisms, especially

Streptomyces or other actinomycetes, transformed with such vector.

This invention also relates to a recombinant DNA molecule comprising the Streptomyces gal operon galE, galT or galk gene, or any functional derivative thereof. By 5 "functional derivative" is meant any derivative of the Streptomyces gal operon galE, galT or galK gene which functions in substantially the same way as the naturally occurring gene in terms of production of an active galE, galT, or galK type gene product. Such 10 derivatives include partial sequences of the Streptomyces gal operon galE, galT, or galK gene, as well as derivatives produced by modification of the gal operon sequence. Techniques for effecting such modification are known in the art, and some have been outlined above. 15 naturally occurring Streptomyces gal operon galE, galT and/or galk gene can be isolated from the naturally occurring Streptomyces gal operon by conventional techniques. The Streptomyces gal operon galE, galT and/or galk gene can be used as a selection marker in an 20 appropriate host mutant. By "appropriate host mutant is meant a host which does not utilize galactose because it contains a defect within a homologous galE, galT and/or galk gene. Thus, a recombinant DNA molecule (comprising the Streptomyces gal operon galE, galT and/or galK gene 25 and a foreign functional DNA sequence, both of which are operatively linked to appropriate regulatory region), which can be prepared by conventional techniques, can be transformed into an appropriate host mutant by conventional techniques for incorporation into the host 30 genome by homologous recombination to enable detection of transformants without the need of maintaining an expensive antibiotic selection. Likewise, a recombinant DNA vector comprising the Streptomyces gal operon galE, galT and/or galk gene and a foreign functional DNA sequence, both of 35 which are operatively linked to appropriate regulatory

1 regions, as well as a replicon, can be transformed into an
appropriate host mutant by conventional techniques to
enable detection of transformants without the need of
maintaining an expensive antibiotic selection. The

Streptomyces gal operon galE, galK and/or galT gene is
also useful for complementation of an appropriate host
mutant.

The Streptomyces gal operon galE gene is also useful for providing a ribosome binding site and 10 initiation codon which can be fused to a foreign functional DNA sequence to enable the expression of such coding sequence when incorporated into an appropriate expression vector and transformed into an appropriate host. If such foreign functional DNA sequence is fused to 15 the galE gene ribosome binding site and initiation codon in a recombinant DNA expression vector comprising the Streptomyces gal operon P2 promoter expression unit, or the entire gal operon, such DNA sequence will be constitutively expressed when such vector is transformed 20 into an appropriate host organism. If such DNA sequence is fused to the galE gene ribosome binding site and initiation codon in a recombinant DNA expression vector comprising the Streptomyces gal operon P2 promoter regulated region, expression of such DNA sequence can be 25 regulated when such vector is transformed into an appropriate host organism by controlling the presence or absence of galactose or glucose.

The Streptomyces gal operon galT gene is also useful for providing a ribosome binding site and 30 initiation codon which can be fused to a foreign functional DNA sequence to enable the expression of such coding sequence when incorporated onto an appropriate expression vector and transformed into an appropriate host. If such DNA sequence is fused to the galT gene 35 ribosome binding site and initiation codon in a recombinant DNA expression vector comprising the

Streptomyces <u>qal</u> operon Pl promoter regulated region, or the entire <u>qal</u> operon, expression of such coding sequence can be regulated in a host transformed with such vector as outlined above.

This invention also relates to a recombinant DNA vector comprising a replicon, Streptomyces gal operon, or a functional and regulatable derivative thereof, and a foreign functional DNA sequence operatively linked to such operon. Such vector can be prepared by conventional techniques. The replicon employed should be one known for its ability to stably and extrachromosomally, maintain a vector in the host organism which is to be the host transformed with the vector.

This invention also relates to a transformed host microorganism comprising a recombinant DNA vector wherein said vector contains a replicon, the Streptomyces gal operon, or a functional and regulatable derivative thereof, and a foreign functional DNA sequence operatively linked to such operon; and to the melhod of preparing such host which comprises transforming an appropriate host microorganism with such vector. Appropriate host microorganisms which may be employed in the method of this invention include viruses, and eukaryotic and prokaryotic cells or organisms, especially actinomycetes, such as those of the genus Streptomyces. The most preferred host microorganisms belong to the genus Streptomyces. Preferred species of Streptomyces include Streptomyces lividans, S. coelicolor, S. azuraeus and S. albus. Transformation of such host microorganism with such vector can be accomplished using conventional techniques such as the method of Chater et al., Curr. Top. Micro. Imm., 96, 69-95 (1982). This invention also related to a method of expressing the functional DNA sequence contained by such transformed host of this invention which comprises cultivating such transformed host under suitable conditions such that the functional DNA sequence is

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1 expressed. By "suitable conditions" is meant those conditions which will allow the host to grow and which enable the expression of the functional DNA sequence. Such suitable conditions can be determined by one of skill 5 in the art using conventional techniques and will depend on various factors, such as the host organism employed and the functional DNA sequence to be expressed. invention is also related to a method of regulating the expression of the functional DNA sequence contained by 10 such transformed host which comprises cultivating a transformed host containing such functional DNA sequence under appropriate conditions such that its expression is regulatable. By "appropriate conditions" is meant those conditions which enable the Streptomyces gal operon (and 15 thus the foreign functional DNA sequence) to be regulatable. By "regulatable" is meant responsive to the presence of galactose or its metabolites and the presence of glucose or its metabolites in the growth media of the transformed host cell. Such regulation can be carried out 20 by addition or deletion of galactose or glucose to the transformed host's culture medium. The optimal levels of galactose and/or glucose for up or down-regulation of the expression of the foreign functional DNA coding sequence by the transformed host of this invention can be readily 25 determined by one of skill in the art using conventional techniques.

This invention also relates to a recombinant DNA vector comprising a replicon, a <u>Streptomyces gal</u> operon P2 promoter expression unit, or a functional derivative thereof, and a foreign functional DNA sequence operatively linked to such unit. Such a vector can be prepared by conventional techniques. The replicon employed should be one known for its ability to stably, and extrachromosomally, maintain a vector in the host organism which is to be transformed with the vector.

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This invention also relates to a transformed host 1 microorganism comprising a recombinant DNA vector wherein said vector contains a replicon, the Streptomyces gal operon P2 promoter expression unit, or a functional derivative thereof, and a foreign functional DNA sequence 5 operatively linked to such unit; and to the method of preparing such host which comprises transforming an appropriate host microorganism with such vector. By the term "operatively linked" is meant that a functional DNA sequence is transcriptionally or translationally linked to 10 an expression control sequence (i.e., the Streptomyces gal operon, P2 promoter expression unit, Pl promoter regulated region, Pl promoter or P2 promoter) in such a way so that the expression of the functional DNA sequence is under control of the expression control sequence. Thus, for 15 example, a foreign functional DNA sequence can be transcriptionally or translationally linked to the Streptomyces gal operon by inserting such operon within the Streptomyces gal operon Pl or P2 promoter transcript. By the term "replicon" is meant that region of DNA on a 20 plasmid which functions to maintain, extrachromosomally, such plasmid in a host microorganism or cell transformed therewith. It has also been discovered that the Streptomyces gal operon, and smaller portions thereof, is useful as a nucleic acid probe to obtain homologous 25 sequences from other cells and organisms. Appropriate host microorganisms which may be employed in the method of this invention include any virus or eukaryotic or prokaryotic cell or organism, especially any actinomycetes 30 such as those of the genus Streptomyces. The most preferred host microorganisms belong to the genus Streptomyces. Preferred species of Streptomyces include Streptomyces lividans, S. coelicolor, S. azuraeus and S. albus. Transformation of such host microorganism with 35 such vector can be accomplished using conventional

techniques such as the method of Chater et al., Curr. Top. Micro. Imm., 96, 69-95 (1982). This invention also related to a method of expressing the functional DNA sequence contained by such transformed host of this invention which comprises cultivating such transformed host under suitable conditions such that the functional DNA sequence is expressed. By "suitable conditions" is meant those conditions which will allow the host to grow and which enable the expression of the functional DNA 10 sequence. Such suitable conditions can be determined by one of skill in the art using conventional techniques and will depend on various factors, such as the host organism employed and the functional DNA sequence to be expressed.

This invention also relates to a recombinant DNA vector comprising a replicon, a Streptomyces gal operon Pl. promoter regulated region, or a functional and regulatable derivative thereof, and a foreign functional DNA sequence operatively linked to such region. Such a vector can be prepared by conventional techniques. The replicon employed should be one known for its ability to stably and extrachromosomally maintain a vector in the host organism which is to be the host transformed with the vector.

This invention also relates to a transformed host microorganism comprising a recombinant DNA vector wherein said vector contains a replicon, a Streptomyces gal operon Pl promoter regulated region, or a functional and regulatable derivative thereof, and a foreign functional DNA sequence operatively linked to such region; and to the method of preparing such host which comprises transforming an appropriate host microorganism with such vector. Appropriate host microorganisms which may be employed include any virus or eukaryotic or prokaryotic cell or organism especially actinomycetes such as those of the genus Streptomyces. The most preferred host microorganisms belong to the genus Streptomyces.

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Preferred species of Streptomyces include Streptomyces lividans, S. coelicolor, S. azuraeus and S. albus. Transformation of such host microorganism with such vector can be accomplished using conventional techniques such as the method of Chater et al., Curr. Top. Micro. Imm., 96, 69-95 (1982). This invention also related to a method of expressing the foreign functional DNA sequence contained by such transformed host of this invention which comprises cultivating such transformed host under suitable conditions such that the functional DNA sequence is expressed. By "suitable conditions" is meant those conditions which will allow the host to grow and which enable the expression of the functional DNA sequence. Such suitable conditions can be determined by one of skill in the art using conventional techniques and will depend on various factors, such as the host organism employed and the functional DNA sequence to be expressed. This invention also related to a method of regulating the expression of the functional DNA sequence contained by such transformed host which comprises cultivating a transformed host containing such functional DNA sequence under appropriate conditions such that its expression is regulatable. By "appropriate conditions" is meant those conditions which enable the Streptomyces gal operon Pl promoter regulated region (and thus the foreign functional DNA sequence) to be regulatable. By "regulatable" is meant responsive to the presence or absence of galactose or its metabolites and the presence or absence of glucose or its metabolites in the growth media of the transformed host cell. Such regulation can be carried out by addition or deletion of galactose or glucose to the transformed host's culture medium.

This invention also relates to a recombinant DNA vector comprising a replicon, a <u>Streptomyces gal</u> operon P2 promoter, or a functional derivative thereof, and a foreign functional DNA sequence operatively linked to such

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promoter. Such a vector can be prepared by conventional techniques. The replicon employed should be one known for its ability to stably and extrachromosomally maintain a vector in the host organism which is to be the host transformed with the vector.

This invention also relates to a transformed host microorganism comprising a recombinant DNA vector wherein said vector contains a replicon, a Streptomyces gal operon P2 promoter, or a functional derivative thereof, and a 10 foreign functional DNA sequence operatively linked to such region; and to the method of preparing such host which comprises transforming an appropriate host microorganism with such vector. Appropriate host microorganisms which may be employed include actinomycetes such as those of the genus Streptomyces. The most preferred host 15 microorganisms belong to the genus Streptomyces. Preferred species of Streptomyces include Streptomyces lividans, S. coelicolor, S. azuraeus and S. albus. Transformation of such host microorganism with such vector can be accomplished using conventional techniques such as 20 the method of Chater et al., Curr. Top. Micro. Imm., 96, 69-95 (1982). This invention also related to a method of expressing the foreign functional DNA sequence contained by such transformed host of this invention which comprises cultivating such transformed host under suitable 25 conditions such that the functional DNA sequence is expressed. By "suitable conditions" is meant those conditions which will allow the host to grow and which enable the expression of the functional DNA sequence. Such suitable conditions can be determined by one of skill 30 in the art using conventional techniques and will depend on various factors, such as the host organism employed and

This invention also relates to a recombinant DNA vector comprising a replicon, <u>Streptomyces gal</u> operon Pl promoter, or any regulatable and functional derivative

the functional DNA sequence to be expressed.

thereof, and a foreign functional DNA sequence operatively linked to such region. Such a vector can be prepared by conventional techniques. The replicon employed should be one known for its ability to stably and extrachromosomally maintain a vector in the host organism which is to be the host transformed with the vector.

This invention also relates to a transformed host microorganism comprising a recombinant DNA vector wherein said vector contains a replicon, the <u>Streptomyces gal</u> operon Pl promoter, or any regulatable and functional derivative thereof, and a foreign functional DNA sequence operatively linked to such region; and to the method of preparing such host which comprises transforming an appropriate host microorganism with such vector.

- 15 Appropriate host microorganisms which may be employed include viruses or prokaryotic or eukaryotic cells or organisms, especially actinomycetes such as those of the genus Streptomyces. The most preferred host microorganisms belong to the genus Streptomyces.
- Preferred species of <u>Streptomyces</u> include <u>Streptomyces</u>

 <u>lividans</u>, <u>S. coelicolor</u>, <u>S. azuraeus</u> and <u>S. albus</u>.

 Transformation of such host microorganism with such vector can be accomplished using conventional techniques such as the method of Chater et al., <u>Curr. Top. Micro. Imm.</u>, <u>96</u>,
- 25 69-95 (1982). This invention also relates to a method of expressing the foreign functional DNA sequence contained by such transformed host of this invention which comprises cultivating such transformed host under suitable conditions such that the functional DNA sequence is
- expressed. By "suitable conditions" is meant those conditions which will allow the host to grow and which enable the expression of the functional DNA sequence. Such suitable conditions can be determined by one of skill in the art using conventional techniques and will depend on various factors, such as the host organism employed and

1 the foreign functional DNA sequence to be expressed. invention also relates to a method of regulating the expression of the functional DNA sequence contained by such transformed host which comprises cultivating a 5 transformed host containing such foreign functional DNA sequence under appropriate conditions such that its expression is regulatable. By "appropriate conditions" is meant those conditions which enable the gal operon Pl promoter (and thus the functional DNA sequence) to be 10 By "regulatable" is meant responsive to the regulatable. presence or absence of galactose or its metabolites and the presence of glucose or its metabolites in the growth media of the transformed host cell. Such regulation can be carried out by addition or deletion of galactose or 15 glucose to the transformed host's culture medium.

EXAMPLES

In the following Examples, specific embodiments of the invention are more fully disclosed. These Examples are intended to be illustrative of the subject invention and should not be construed as limiting its scope. In all Examples, temperature is in degrees Centigrade (°C).

By utilizing conventional methods, such as those outlined in the following Examples, one of skill in the art can isolate the gal operon from any galactose utilizing strain of Streptomyces. Furthermore, by utilizing techniques similar to those employed herein to isolate the Streptomyces gal operon, one of skill in the art can attempt to use the Streptomyces gal operon to isolate a gal operon from other galactose utilizing other strains of Streptomyces, especially S. coelicolor, S. azuraeus, S. albus and other S. lividans strains.

Molecular genetic manipulations and other techniques employed in the following Examples are described in Hopwood et al., <u>Genetic Manipulation of Streptomyces: A Laboratory Manual</u>, John Innes Foundation, Norwich, England (1985).

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                                 ABBREVIATIONS
                In the following Examples, the following
      abbreviations may be employed:
                      10 grams (g) tryptone, 5 g yeast extract, 5g
                LB:
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     NaC1
                MBSM (modified MBSM): See, Brawner et al., Gene,
      40, 191 (1985) (in press)
      MOPS: (3)-N-morpholino-(proprane-sulfonic acid)
10
                YEME + MgCl<sub>2</sub> + Glycine: [per liter(1)] 3 g
      yeast extract, 5 g peptone, 3 g malt extract, 10 g
      glucose, 10 g {\rm MgCl}_2 62{\rm H}_2{\rm O}, 340 g sucrose.
                SL: Mix together (NH_4)_2SO_4(1g/1);
      L-asparagine (2 g/l); K_2^{HPO}_4 (9 g/l); NaH_2^{PO}_4 (1
15
      g/l) for 0.2% agar and autoclave. Then mix with yeast
      extract (20 g/1), MgCl_2 (5 g/1); CuCl_2 (0.1 g/1);
      Trace elements [20 ml/l - include ZnCl2-40 mg/l;
      FeCl<sub>3</sub>"6H<sub>2</sub>O (200 mg/l); CuCl<sub>2</sub>"2H<sub>2</sub>O (10 mg/l);
      NaB_4O_7 10H<sub>2</sub>O (10 mg/1); (NH_4)_6MO_7O_{24} 4
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      H_2O(10 \text{ mg/l}) filter and sterilize.
                YEME (Ym base): (per liter) yeast extract (3g);
      peptone (5g); malt extract (3g); MgCl<sub>2</sub>*6H<sub>2</sub>O (2g)
                 Ymglu: YEME + glucose (10g)
                 Ymgal: YEME + galactose (10g)
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BACTERIAL STRAINS

In the following Examples, the following strains of \underline{E} . \underline{coli} are employed:

	CGSC Strain #(a)	Strain Designation	Sex	Chromosomal Markers
	4473 (<u>gal</u> E ⁻)	W3109		galE9, (b) g-; IN(rrnD-rrnE)1
	4467 (<u>gal</u> T ⁻)	W3101	F ⁻	galT22 ^(b) g ⁻ ;IN(rrnD-rrnE)l
10	4498 (<u>gal</u> E ⁻)	PL-2	Hfr	thi-1, relAl, 921E28,g, spoT1

(a) CGSC Strain # is the stock number designated for such strain by the E. coli Genetic Stock Center of the Department of Human Genetics, Yale University School of Medicine, 333 Cedar Street, P.O. Box 3333, New Haven, Connecticut, 06510, U.S.A.
(b) galE9 is the old Lederberg gal9; galT22 is the old Lederberg gal1.

20 <u>S1 ANALYSIS</u>

Sl analysis is used to identify the 5' end of RNAs and the length of a RNA of interest. In the following Examples, Sl analysis refers to Sl experiments carried out according to the method of Weaver et al., Nucl. Acids Res., 7, 1175 (1979) and Berk et al., Proc. Natl. Acad. Sci. USA, 75, 1214 (1978).

EXAMPLE I

A. CLONING OF A STREPTOMYCES LIVIDANS GALACTOKINASE GENE.

Streptomyces lividans strain 1326 is described by Bibb et al., Mol. Gen. Genetics, 184, 230-240 (1981) and was obtained from D. A. Hopwood, John Innes Foundation, Norwich, England. Streptomyces lividans strain 1326 and S. lividans strain 1326 containing the pIJ6 plasmid were deposited in the Agricultural Research Culture Collection,

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Peoria, Illinois, U.S.A., on June 1, 1982, under accession numbers NRRL 15091 and 15092, respectively.

High molecular weight chromosomal DNA was isolated from <u>Streptomyces lividans</u> strain 1326 according to the method of Maniatis et al., "Molecular Cloning. A Laboratory Manual", Cold Spring Harbor Laboratory (1982) and was size fractionated on a 10-40% sucrose gradient (See, Maniatis et al., cited above, p. 284-285).

10 Fractions of 18-24 kilobase (Kb) pairs were combined and dialyzed exhaustively against 10 mM Tris-HCl/l mM EDTA (pH 8). Cosmid shuttle vector pJW357 was employed to clone such fractionated chromosomal DNA in its entirety. pJW357 was constructed by fusing pDPT6 cut with PstI to pIJ350

15 cut with PstI. pIJ350 is described in Kieser et al., Mol. Gen. Genet., 185, 223-238 (1982). pDPT6 is a tetracycline and chloramphenical resistant, pBR322-based E. coli cosmid cloning vector described in Taylor et al., U.S. Patent No. 4,476,227. pJW357 has a unique EcoRI site in the chloramphenical resistance gene and a unique BamHI site in

the Tc^R (tetracycline) resistance gene. pJW357 was digested with BamHI, dephosphorylated with alkaline phosphatase, and ligated to the fractionated chromosomal DNA described above.

The ligation product was packaged into bacteriophage heads (using the in vitro packaging system described by Maniatis et al., cited above, p. 264-265) and transfected into E. coli strain K21 which is a galk derivative of E. coli MM294. The transformation culture was grown for two hours in LB and for an additional two hours in LB with 25 ug/ml chloramphenicol, washed three times with equal volumes of M9 media [see, Miller, "Experiments in Molecular Genetics", Cold Spring Harbor Laboratory (1972)] without a carbon source, and plated onto M9 agar [supplemented with proline, histidine, arginine, isoleucine, leucine, saline and .5% galactose;

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See, Adams et al., Biochem. Biophys. Res. Comm., 89(2), 650-58 (1979)] with 30 mg/ml chloramphenicol. Twenty plates were spread with approximately 200 transformants per plate. After three days incubation at 37°C, no transformants were detected. The minimal plates were then sprayed with nicotinic acid to 5 ug/ml to supplement the nicotinic acid requirement of E. coli strain K21, and the incubation was continued for 3 more days at 37°C and for 2 additional days at room temperature. After such incubation, the surviving colonies were patched to both MacConkey galactose agar (MAC-GAL) [See, Miller et al., cited above] with 30 ug/ml chloramphenicol and to M63 minimal agar [See, Miller et al., cited above] supplemented with .5% galactose, 5 ug/ml nicotinic acid, 5 ug/ml thiamine and 30 ug/ml chloramphenicol. Only two colonies contained cosmid DNA that transformed E. coli K21 to a galk phenotype. Such cosmids were designated as pSLIVGAL-1 and pSLIVGAL-2. Both colonies were light red on MAC-GAL (i.e., they were $gal K^+$) and also grew on the M63 medium.

Plasmids pSLIVGAL-1 and pSLIVGAL-2 were isolated from the two galk colonies described above and were transformed, according to the method of Chater et al.,

Curr. Top. Micro. Imm., 96, 69-95 (1982), into Streptomyces

lividans strain 1326-12K (a galk deficient strain isolated after UV mutagenesis of S. lividans strain 1326, See,

Brawner et al., Gene, 40, 191 (1985), (in press). Plasmid encoded complementation of the S. lividans 1326-12K

(galk) host was tested by observing growth of spores plated on MBSM-gal-thiostrepton according to the method of Brawner et al., Gene, 40, 191 (1985) (in press).

pSLIVGAL-2 showed no detectable complementation of the Streptomyces 1326-12K host.

Cell extracts were prepared from cultures grown
in SL medium supplemented with 1% glucose or galactose and
10)g/ml thiostrepton. The extracts were analyzed for

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galactokinase production by immunoblot analysis (see, Brawner et al., <u>Gene</u>, <u>40</u>, 191 (1985), in press) using rabbit antisera prepared against <u>E. coli</u> galactokinase. The protein detected by immunoblot analysis was the approximate size of <u>E. coli</u> galK. Such protein appeared in galactose supplemented cultures of <u>Streptomyces</u> at levels several fold higher than in glucose cultures.

10 B. MAPPING OF THE S. LIVIDANS GALK REGION WITHIN A COSMID.

The galk region of the pSLIVGAL1 and pSLIVGAL2 cosmids, prepared as described above, was identified by cloning random fragments from the cosmids into a pUC18 derivative [See, Norrander et al., Gene, 26, 101-106 (1983)] and scoring complementation of E. coli strain MM294 (galk) on MAC-GAL medium. The cosmid clone was partially digested with Sau3AI (using conditions which maximized the yield of 2 to 4 kilobase fragments), and the products of this reaction were ligated into the BglII site of pUC18-TT6, a derivative of pUC18 constructed by insertion of the following synthetic DNA sequence into the BamHI site of pUC18:

5'GATCAGATCTTGATCACTAGCTAGCTAG 3'

TCTAGAACTAGTGATCGATCGATCCTAG 5'

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Twelve galK⁺ clones (red on MAC-GAL) were screened for size. One clone, designated as plasmid pSAUlO, was the smallest and had an insert size of approximately 1.4 Kb.

In contrast to colonies containing pSLIVGAL1, the pUC clones were very red on MAC-GAL medium, indicating an increased production of galactokinase. The most likely explanation for the increased enzyme level was that the <u>S. lividans gal</u>K gene was now being transcribed by an <u>E. colipromoter</u> which was stronger than the upstream promoter on the cosmid.

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The insert of pSAUlO was isolated as an EcoRI to HindIII fragment (these sites flank the insert region of pUC18-TT6) for use as a probe for the S. lividans galk The chromosomal DNA used in the cloning was restricted with EcoRI plus MluI and BamHI plus BqlII, and then blotted according to the method of Southern, J. Mol. Biol., 98, 503 (1975). The pSAU10 fragment was nick translated and hybridized to the blot. The probe identified a 1.3 kb EcoRI-MluI fragment and a 5 kb BamHI-BglII fragment in the chromosomal digests. When this data was compared to the map of the cosmid insert, the location of the galK gene (between map positions 5 and 7, See Table A) was confirmed.

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C.

DNA SEQUENCING OF THE S. LIVIDANS GAL OPERON. The Streptomyces lividans gal operon was sequenced by chain termination [(See, Sanger et al., Proc. Nat'l Acad. Sci., U.S.A., 74, 5463 (1977)] and chemical cleavage [See, Maxam and Gilbert, Methods in Enzymology, 20 65, 499 (1980)]. The initial sequences of galk were derived from Sau3AI and SalI fragments of the insert of pSAU6 (a 2.3 Kb sibling of pSAU10) shotgun cloned into the BamHI and SalI sites (respectively) of M13 mp 10 [See, Messing, Methods in Enzymology, 101, 20 (1983)]. Amino 25 acid sequences of the S. lividans galT, galE and galK genes were predicted by computer, and further analyzed by comparison with amino acid sequences of the E. coli and or S. cerevisiae galactokinase, gal-l-phosphate uridyltransferase and UDP-4-epimerase enzymes. 30 sequences of these proteins were predicted by computer analysis using the total or partial DNA sequence of the genes which encode the gal enzymes [see, Debouck et al., Nuc Acids. Res., 13(6), 1841-1853 (1985), and Citron and Donelson, J. Bacteriology, 158, 269 (1984)]. 35 homology was found between the inferred protein sequence

for the \underline{S} . lividans galk, galt, galt gene products and their respective \underline{E} . coli and/or \underline{S} . cerevisiae gene products.

The complete DNA sequence of the <u>S. lividans gal</u> operon is shown in Table 1. Included in Table 1 are the transcription start sites for the operon's promoters and the predicted amino acid sequences of the <u>galT</u>, <u>galE</u> and <u>galK</u> gene products.

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TABLE 1 TRANSLATED SEQUENCE OF STREPTOMYCES LIVIDANS GALACTOSE OPERON

-120 -110 -100 -90 -80 -70.~ CTA CGC CTC CGC GTT CAG TAA TTG AAC ACT TTT GGT GAT GAA CTT TGT TTG ATT GTG 10 -60 -50 -40 -30 -20 galP1 -10 10 20 40 15 ACG GGC GTC CTG GTG ACT CAT GGG TGG GTG CAG AGG AGT GCG GCA GTG AAG AAC ACC Wet Thr His Gly Trp Val Cln Arg Ser Ala Ala Val Lys Lys Thr galT 70 50 60 80 90 100 TCG ACC CGG CTG GCC GAC GGC CGT GAG CTG GTC TAC TAC GAC CTG CGC GAC GAC ACC Ser Thr Arg Leu Ala Asp Gly Arg Glu Leu Val Tyr Tyr Asp Leu Arg Asp Asp Thr 20 110 120 130 140 GTG CGC GAC GCC GTG GAC CGC CGT CCG CTG GAG CGG ACC GTC ACC ACG TCC GAG GTG Val Arg Asp Ala Val Asp Arg Arg Pro Leu Glu Arg Thr Val Thr Thr Ser Glu Val 160 170 180 190 200 210 25 CGA CGC GAC CCG CTG CTC GGC GAC TCC GCG CCG TCG CGC CTC GCA CCG GCA GGG GCG Arg Arg Asp Pro Leu Leu Gly Asp Ser Ala Pro Ser Arg Leu Ala Pro Ala Gly Ala 220 230 240 250 CAC CTA CCA TCC GCC GGC CGA CCA GTG CCC GCT GTG CCc GTC GGA CGG GGA ACG GCT His Leu Pro Ser Ala Gly Arg Pro Val Pro Ala Val Pro Val Gly Arg Gly Thr Ala 30 280 290 300 310 320 330

GAG CGA GAT CCG GCC TAT GAC GTG GTG GTC TTC GAG AAT CGC TTT CCC TCG CTG GCC Glu Arg Asp Pro Ala Tyr Asp Val Val Phe Glu Asn Arg Phe Pro Ser Leu Ala

l 'able 1 - (cont'd)

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	340						350			360)		3	70			380		
5	GGT Gly	GAC Asp	TCC Ser	GGG Gly	CGC Arg	TGC Cys	GAG Glu	GTC Val	GTC Val	TGC Cys	TTC Phe	ACC Thr	TCC Ser	GAC Asp	CAC His	GAC Asp	GCC Ala	TCC Ser	TTC Phe
	390)		4	100			410			420)		4	130			440	
	GCC Ala	GAC Asp	CTG Leu	AGC Ser	GAG Glu	GAC Glu	CAG Gln	GCC Ala	CGG Arg	CTG Leu	GTC Val	GTC Val	GAC Asp	GCC Ala	TGG Trp	ACG Thr	GAC Asp	CGC Arg	ACC Thr
10		450	0		4	60			470			480)		4	190			500
	TCC Ser	GAG Glu	CTG Leu	TCC Ser	CAT His	CTG Leu	CCC Pro	TCC Ser	GTT Val	GAA Glu	CAG Gln	GTC Val	TTC Phe	TGC Cys	TTC Phe	GAG Glu	AAC <u>Asn</u>	CGG Arg	GJ X
			51	0		ţ	520			530			540)			550		
15	GCC Ala	GAG Glu	ATC Ile	GGG Gly	GTG Val	ACG Thr	CTG Leu	GGT Gly	CAC Bis	CCG Pro	CAC	GCC Gly	CAG Gln	ATC Ile	TAC Tyr	GCC	TAC Tyr	CCG Pro	TTC Phe
	560				0		;	580			590			60	0		(510	
	ACC Thr	ACC	CCC Pro	CGC Arg	ACC Thr	GCC Ala	CTG Leu	ATG Met	CTC Leu	CGT Arg	TCA Ser	CTC Leu	GCC Ala	GCC Ala	CAC His	AAG Lys	GAC Asp	GCG Ala	ACG Thr
20		620)		63	0			640			650			66	0			670
	GGC Gly	GGC Gly	GGG Gly	AAC Asn	CTG Leu	TTC Phe	GAC Asp	TCC Ser	GTG Val	CTG Leu	GAG Glu	GAG Glu	GAG Glu	CTG Lev	GCC	GGT Gly	GAG Glu	CGG Arg	GTC Val
			680)		69	o			700			710			72	0		
25	GTC Val	CTC Lev	G GAG	GGT Gly	GAG Glu	CAC His	TGG Trp	GCC	GCC	TTC Phe	CTC Val	GCG Ala	TAC Tyr	GGC Gly	GCG	CAC His	TGG Trp	CCG Pro	TAC Tyr
	730			740)		75	0			760			770)		78	0	
	GAC Glu	GTC	CAC His	CTC Lev	TAC Tyr	CCG Pro	AAG Lys	CGG Arg	CGG Arg	CTC Val	Pro	GAT Asp	CTG Leu	CTC Leu	GGG Gly	CTC Leu	GAC Asp	GAG Glu	GCG Ala
30		790			800)		81	0			820			830			84	0
	GCT Ala	CGC A Ar	C ACA	A GAM	TTC Phe	CCC Pro	AAG Lys	GTC Val	TAC Tyr	CTC Let	GAG Glu	CTG Leu	CTG Leu	AGG	CGT Arg	TTC Phe	GAC Asp	CGG Arg	ATC Ile

able 1 - (cont'd) -37-TTC GGC GAG GGC GAG CCC CCG ACC CCC TAC ATC GCG GCC TGG CAC CAG GCG CCG TTC Phe Gly Glu Gly Glu Pro Pro Thr Pro Tyr Ile Ala Ala Trp His Gln Ala Pro Phe 🗍 GGG CAG CTG GAG TTC GAG GGT GTG ACG CGC GAC GAC TTC GCG CTC CAC CTG GAA CTT Gly Gln Leu Glu Phe Glu Gly Val Thr Arg Asp Asp Phe Ala Leu His Leu Glu Leu TTC ACT TCC GCC GTA CCT CCG GCA AGC TGA AGT TCC TCG CGG GCT CCG AAT CCG GCA Phe Thr Ser Ala Val Arg Pro Ala Ser --galP2 TGAACG TGTTCATCAA CGACGTACCC CCGGAGCGCG CGGCCGAGCG ACTGCGAGAG GTAGCGAG TTC ATG AGC GGG AAG TAC CTG GTG ACA GGT GGT GCC GGA TAC GTC GGC AGC GTC GTC Met Ser Gly Lys Tyr Leu Val Thr Gly Gly Ala Gly Tyr Val Gly Ser Val Val galE GCC CAG CAC TTG GTG GAG GCG GGG AAC GAG GTC GTG GTG CTG CAC AAT CTG TCG ACC Ala Gln His Leu Val Glu Ala Gly Asn Glu Val Val Val Leu His Asn Leu Ser Thr GGC TTC CGT GAG GTG TGC CGG CGG GTG CCT CGT TCG TCG AGG CGA CAT CCG GGA CGC Gly Phe Arg Glu Val Cys Arg Arg Val Pro Arg Ser Ser Arg Arg His Pro Gly Arg CGC CAA GTG CGT GGA CGG CTC TCG TTC GAC.GGC GTG CTG CAC TTC GCC GCC TTC TCC Arg Gln Val Arg Gly Arg Leu Ser Phe Asp Gly Val Leu His Phe Ala Ala Phe Ser CAG GTC GGC GAG TCG GTC GTG AAG CCC GAG AAG TAC TGG GAC AAC AAC GTC GGT GGC Gln Val Gly Glu Ser Val Val Lys Pro Glu Lys Tyr Trp Asp Asn Asn Val Gly Gly

-38-Tabel 1 - (cont'd) Thr Wet Ala Leu Leu Glu Ala Wet Arg Gly Ala Gly Val Arg Arg Leu Val Phe Ser TCC ACG GCC GCC ACG TAC GGC GAG CCC GAG CAG GTT CCC ATC GTC GAG TCC GCG CCG Ser Thr Ala Ala Thr Tyr Gly Glu Pro Glu Gln Val Pro Ile Val Glu Ser Ala Pro ACG AGG CCC ACC AAT CCG TAC GGC GCC TCG AAG CTC GCC GTC GAC CAC ATG ATC ACC Thr Arg Pro Thr Asn Pro Tyr Gly Ala Ser Lys Leu Ala Val Asp His Wet Ile Thr GGC GAG GCG GCC CAC GGG CTG GGC GCG GTC TCC GTG CCG TAC TTC AAC GTC GCG Gly Glu Ala Ala Ala His Gly Leu Gly Ala Val Ser Val Pro Tyr Phe Asn Val Ala GGC GCG TAC GGG GAG TAC GGC GAG CGC CAC GAC CCC GAG TCG CAT CTG ATT CCG CTG Gly Ala Tyr Gly Glu Tyr Gly Glu Arg His Asp Pro Glu Ser His Leu Ile Pro Leu GTC CTT CAA GTG GCG CAG GGC AGG CGG GAG GCC ATC TCC GTC TAC GGC GAC GAC TAC Val Leu Gln Val Ala Gln Gly Arg Arg Glu Ala Ile Ser Val Tyr Gly Asp Asp Tyr CCG ACG CCG GAC CGA CCT GTG TGC GCG ACT ACA TCC ACG TCG CCG ACC TGG CCG AGG Pro Thr Pro Asp Arg Pro Val Cys Ala Thr Thr Ser Thr Ser Pro Thr Trp Pro Arg CCC ACC TGC TGG CCG TGC GCC GCC CCG GGC GAG CAC CTC ATC TGC AAC CTG GGC Pro Thr Cys Trp Pro Cys Ala Ala Ala Pro Gly Glu His Leu Ile Cys Asn Leu Gly

AAC GGC AAC GGC TTC TCC GTC CGC GAG GTC GTC GAG ACC GTG CGG CGG GTG ACG GGC Asa Gly Asa Gly Phe Ser Val Arg Glu Val Val Glu Thr Val Arg Arg Val Thr Gly

1	Cable 1	. - (cont	' d)							-	39-							
	1	880			1890			190	0		1	910			1920	•		193	0
5																GCG Ala			GTC Val
		11	940			1950			196	0		1	970			1980			1990
																			CTC Leu
10			20	000		:	2010			202	0		2	030		:	2040		
																CAG Gln		TA	
		20	050		:	2060			2070)		2	080			2090			2100
15	ACC	GCA	GTT	ACC	GGA	AAG	GCG	ACC	GGT	CAG	GGC		Gly			GTC Val			
				110			2120			2130		Ü	2	140			2150		
20	TCG Ser	CCC Ala	AGC Ser	GGT Gly	TCC Ser	GGG Gly	AGC Ser	TGT Cys	ACG Thr	GGG Gly	CGG Arg	AGC Ser	CGG Arg	AGG Arg	GGG Gly	TGT Cys	GGG Gly	CGC Arg	CGA Arg
_ •	2160				70			2180			2190			_	200			2210	
	<u>Ala</u>	GGC Gly	CGC Arg	GAG Glu	AAC Asn	CTC Leu	ATC Ile	GGG Gly	GAG Glu	CAC His	ACC Thr	GAC Asp	TAC Tyr	AAC Asn	GAC Asp	GGC Gly	TTC Phe	GTC Val	ATG <u>We</u> t
2.5		2220)		22	230		2	2240			2250)		23	260		2	270
25	CCT Pro	TCG Ser	CCC Pro	TGC Cys	CGC Arg	ACC Thr	AGG Arg	TCG Ser	CGC Arg	CCG Pro	TCT Ser	CCC Pro	GGC Gly	GCG Ala	AAC Asn	GAC Asp	GGC Gly	ATC Ile	CTG Leu
			2280)		22	290		2	300			2310)		23	20		
30	CGC Arg	CTG Leu	CAC	TCG Ser	GCC Ala	GAC Asp	GTC Val	GAC Asp	GCC Ala	GAC Asp	CCG Pro	GTC Val	GAG Glu	CTG Leu	CGC Arg	CTC Val	GCC Ala	GAC Asp	CTG Leu
	2330			2340)		23	50		2	360			2370)		23	80	
	GCC Ala	CCC Pro	GCG Ala	TCG Ser	GAC Asp	AAG Lys	TCC Ser	TGG Trp	ACG Thr	GCG Ala	TAC Tyr	CCC Pro	TCG Ser	GGC Gly	GTC Val	CTG Leu	TGG Trp	GCG Ala	CTG Leu

l table 1 - (cont'd)

-40-

	2390		2400		2410		2420		2430		2440
5	CGC GAG (GCC GGA	CAC GAG His Glu	CTG AC	cc GGC	GCC G	AC GTC	CAC CTG	GCC TCG Ala Ser	ACC GT Thr Va	c ccc
	2	450	2450)	24	70	2	2480	249	0	
	TCC GGG Ser <u>Gly</u>	GCG GGG	CTC TCC Leu Ser	TCC TC Ser Se	CC GCG	GCC C	CTG GAG Leu Glu	CTC CGT Val Arg	CCC CTG	GCG AT Ala Me	G AAC t Asn
10	2500	2510		2520		253	30	2540)	2550	
	GAC CTG Asp Leu	TAC GCC Tyr Ala	CTC GCG Leu Ala	CTG CC	GC GGC	TGG (CAG CTG Gln Leu	GCC CGC	CTG TGC Leu Cys	CAG CG Gln Ar	C GCG
	2560		2570	25	580		2590		2600	26	10
15	GAG AAC Glu Asn	GTC TAC	GTC GGC Val Gly	GCC CC	CC GTC	GGC A	ATC ATG Ile Wet	GAC CAC	ACG GCC	TCC GC	C TGC
	26	520	2630		2640)	2	650	2660)	2670
	TGC GAG Cys Glu	GCG GGC	ACG CCC Thr Pro	TCT TO	CC TCG	ACA (Thr l	CCC GCG Pro Ala	ACC TC	CCC AGO	GGC AC	A TCC
20		2680		2690		2700		2710		2720	
	CCT TCG Pro Ser	ACC TCG Thr Ser	CCG CCG Pro Pro	AGG G	CA TGC	GCC C	TGC TGG Cys Trp	TCG TC	G ACA CCC	GGG TO	CA AGC er Ser
	2730	2	740	27	50	:	2760		2770	278	30
25	ACT CCC Thr Pro	ACA GCG Thr Ala	AGG GCG	AGT A Ser T	CG GCA	AGC Ser	Ala Ala	CGG GC	T GCG AGA	AGG GG Arg Al	CG CCG la Pro
	279	0	2800		2310		282	20	2830		2840
	CGC TGC Arg Cys	TGG GCC	TCG ACC	CGC T	GC GAC Cys Asp	GTG Val	CCG TAC	GCC GA	C CTG GA	C GCG GG P Ala A	CG CTG la Leu
30		2850	2	2860		2870		2880	:	2890	
	GAG CGG Glu Arg	CTG GGG	GAC GAC	GAG G	GAG GTG Glu Val	CGC Arg	CGC CTC	GTC CG	G CAC GT g His Va	G GTG A	CC GAG hr Glu

Table 1 - (cont'd) -41-GAC GAG CGC GTC GAA CGG GTG GTC GCG CTG CTG GAG TCG GCG ACA CCC GGC GCA TCG Asp Glu Arg Val Glu Arg Val Val Ala Leu Leu Glu Ser Ala Thr Pro Gly Ala Ser GCG CCG TCC TGG TCG AGG GCC ACG CCT GCT GCG CGA CGA CTT CCG CAT CTC CTG CCC Ala Pro Ser Trp Ser Arg Ala Thr Pro Ala Ala Arg Arg Leu Pro His Leu Leu Pro CGA GCT GGA CCT GGT CGA CAC GGC CCT GGC CTC CGC GGC CCT CGG CGC CGG ATG Arg Ala Gly Pro Gly Arg Arg His Gly Pro Gly Leu Arg Gly Pro Arg Arg Met ACC GGC GGC GGC TTC GGC GGC TCG GCG ATC GTC CTG GTG GAG GCC GCC GCG GTG GAC Thr Cly Cly Cly Phe Cly Cly Ser Ala Ile Val Leu Val Clu Ala Ala Ala Val Asp GCC GTC ACC AAG GCG GTC GAG GAC GCC TTC GCC GCG GCG GGC CTC AAG CGT CCG CGG Ala Val Thr Lys Ala Val Glu Asp Ala Phe Ala Ala Ala Gly Leu Lys Arg Pro Arg GTG TTC GAG GCG GTG CCT CGG CGG GGC GCG CCT GGT CTG ACG GTC AGC CGA GCC Val Phe Glu Ala Val Pro Arg Arg Gly Ala Ala Pro Gly Leu Thr Val Ser Arg Ala GCT TCA CCA GCG TGT ACT CCG TGA TCC CCG GCG GGT AGT CGG GGA TCA CGC ACA TGA Ala Ser Pro Ala Cys Thr Pro ---GCT GCT AGC CGC

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EXAMPLE 2

PROMOTERS OF THE S. LIVIDANS GAL OPERON

a) Pl promoter

(i) Summary

This promoter is galactose inducible, glucose repressible and is the regulatable promoter for the entire <u>Streptomyces gal</u> operon. Sl data indicates that the <u>Streptomyces lividans gal</u> operon encodes a polycistronic transcript of approximately 3.4 kilobases

(Kb). The transcript consists of approximately 1 Kb for galT, followed by approximately 1 Kb each for galE and galK. (See, Figure 1).

Galactose induction of Pl is mediated, at least in part, by an operator sequence whose 5' end is located 31 bp upstream of the transcription start site and a repressor protein which recognizes the operator.

(ii) Experimental: Isolation, Localization, and Characterization of the Pl promoter.

20 The sequences upstream of the Streptomyces lividans galk ATG were screened for promoters using the E. coli galk promoter probe system of Brawner, et al., Gere, 40, 191, (1985), in press. The HindIII-MluI fragment (See, Table A, map positions 1-5) was restricted with 25 Sau3AI, ligated into the unique BamHI site of pK21 (Figure 2), and transformed into E. coli K2l (galK) according to the method of Example 1. pK21 is a derivative of pSKO3 and is an E. coli-Streptomyces shuttle vector containing the E. coli galk gene (See, Figure 2). The construction 30 of pSKO3 is described in Rosenberg et al., Genetic Engineering, 8, (1986), in press. The clones which expressed galk, i.e., those which had promoter activity, were identified on MacConkey - galactose plates. galk clones (designated as pK21 MH1 and 2) were 35 transformed into Streptomyces 1326-12K (galK).

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Extracts from transformants were cultured in Ymglu and Ymgal, and were analyzed by western blot analysis using anti-E. coli galactokinase antiserum. The blots showed significantly higher levels of galactokinase in the extracts from the galactose induced cultures.

pK21 MH1 and 2 were shown by restriction analysis to contain a 410 bp Sau3AI insert which is contained within the HindIII and Bq1II sites (see Table A, map positions 1-2) by Southern blot analysis according to the method of Southern, J. Mol. Biol., 98, 503 (1975). The cloned fragment was analyzed by S1 analysis using RNA isolated from Streptomyces lividans 1326-12K and E. coli K21 cultures. The fragment yielded a 290 nucleotide protected fragment after S1 digestion (indicating the 5' end of an mRNA 290 bp upstream of the Sau3AI site). Hybridization experiments (using single stranded M13 clones of this region) have identified the direction of transcription as left to right as shown in Figure 2 (i.e., transcription is going toward galK).

Conventional DNA sequence analysis and additional S1 mapping analysis were used to define the 5 end of the mRNA.

The sequences responsible for regulating galactose induction of Pl were localized by removing sequences upstream of the transcription start site by nuclease Bal31. Any change in promoter function or galactose induction by removal of these sequences was assessed using the \underline{E} . \underline{coli} \underline{galK} promoter probe plasmid used to identify Pl.

(iii) Construction of Gal Promoter Deletions.

Plasmid pHL5 was constructed by cloning a DNA fragment containing 100 bp of sequences downstream from the start of Pl transcription and 216 bp upstream from the start of Pl transcription into plasmid pUCl9TTl. Plasmid pUCl9TTl is described in Norrander et al., Gene, 26, 101-106 (1983) and has the Unker as pUCl8-TT6. See, Example IB. Deletions extending into the upstream

- l sequence preceeding Pl were generated by linearizing pHL5 with HindIII and treating the ends with nuclease Bal31.

 The uneven ends were subsequently repaired with the Klenow fragment of DNA polymerase I. Bal31-treated pHL5 was then
- fragments in the molecular weight range of 100-300 bp were eluted from the gel and subcloned into M13 mp 10 that had been digested with HindII and BamHI. [See, Messing, Methods in Enzymology, 101, 20 (1983)]. Individual
- 10 deletions were then sequenced from the single stranded phage DNA the dideoxy chain termination method of Sanger, et al., cited above.
 - (iv) Linking the Pl Promoter Deletions to the E. coli galk Gene.
- The various mp 10 clones were digested with BamHI
 and HindIII. DNA fragments containing individual deletions were isolated from low-melting point agarose gels and then ligated to pK21 (see, Figure 2) that had been digested with BamHI and HindIII. After
- 20 transformation into E. coli MM294, plasmid DNA was isolated for each of the deletion derivatives and transformed into Streptomyces Lividans 12K.
 - (v) Functional Assessment of Bal

31-Generated Deletions in S. lividans

- 25 For each individual promoter deletion, a single thiostrepton resistant transformant was grown to late log in YM base (YEME) + 10 ug/ml thiostrepton. Cells were then pelleted, washed once in M56 media and resuspended in M56 media (see Miller, et al., cited above). The washed
- 30 cells were then used to inoculate YM + 0 1M MOPS (pH 7.2) + 10 ug/ml thiostrepton supplemented with 1% galactose or 1% glucose. The cells were grown for 16 hours then assayed for galactokinase activity.

Ten individual pK2l derivatives containing either 35 120, 67, 55, 34, 31, 24, 20, 18, 10 or 8 bp of sequence upstream of the Pl transcription start site were analyzed

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for galactokinase expression. These results showed that all the information necessary for galactose induction of Pl, (i.e., 10-20 fold greater levels of galactokinase produced in galactose grown cells versus glucose grown cells) is included in the 31 bp of sequence upstream of Pl. A deletion which leaves 34 bp of sequence upstream of Pl is partially inducible by galactose since galactose induced 6-fold greater amounts of galactokinase. one end of the operator must be situated within the sequences between the -24 and -31 position. The remaining deletions which leave either 20, 18, 10 or 8 bp of upstream sequence result in a constitutive Pl promoter, that is the levels of galactokinase produced were equivalent when cells were grown in the presence of galactose or glucose. Although the promoter deletions which retained 8 and 10 bp of Pl were constitutive, the amount of galactokinase produced was reduced 10 fold in comparison to the promoter deletions which retained 18 to 120 bp of upstream sequence. This result indicates that sequences between the -10 and -18 positions of -1 are essential for promoter function.

This data supports a model in which galactose induction of Pl is mediated, at least in part, by an operator sequence. One end of this sequence is 24 to 31 bp upstream of the Pl transcription start site. Removing part or all of the operator results in a promoter which is partially or totally derepressed. The other end of this sequence has not been defined by these experiments but it most likely is contained within the 24 to 31 bp of sequence upstream of the Pl transcription start site. In addition we cannot eliminate the possibility that the 3' end of the operator is also within the 100 bp downstream of the transcription start site since these sequences were contained within the smallest region needed to achieve galactose induction. These data also suggest that the factor which interacts with the operator sequence is a

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repressor protein. Finally, we do not have any evidence which eliminates the possibility that Pl may be controlled by factors other than a repressor (i.e., positive activator such as lambda phage cII protein) to modulate galactose induction promoter transcript.

b) P2 promoter

(i) Summary

The P2 promoter of the <u>Streptomyces gal</u>

10 operon is upstream of the <u>gal</u>E gene and transcribes both galE and <u>gal</u>K genes.

P2 promoter expression is constitutive (i.e., not glucose repressed/galactose induced) as shown by S1 analysis.

(ii) Experimental: Isolation, Localization, and Characterization of the P2 promoter.

The existence of the <u>Streptomyces</u> gal operon P2 promoter became apparent when the <u>BqlII-MluI</u> fragment (see, Table A, map positions 2-5) of <u>S. lividans</u> 1326 DNA was inserted into plasmid pK2l (see, Figure 2) and galactokinase expression was observed in <u>Streptomyces</u> lividans 1326-12K transformed therewith.

DNA sequence analysis and S1 analysis were used to identify the 5' end of the \underline{S} . <u>lividans gal</u> operon P2. The 5' end of the P2 promoter transcript is within 100 bp upstream of the predicted <u>gale</u> ATG.

EXAMPLE 3

EVIDENCE OF A POLYCISTRONIC MESSAGE IN THE STREPTOMYCES GAL OPERON

Sl analysis was used to map the transcripts upstream and downstream of the <u>Streptomyces lividans gal</u> operon <u>gal</u>K gene. In general, overlapping DNA fragments of 1-2 Kb were isolated from subclones, further restricted, and end labelled. The message was followed from the 3' end of <u>gal</u>K to the upstream end at Pl.

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operon transcript probably occurs within the first hundred bases downstream of galk. Fragments 3' labelled at sites within the galk sequence were not protected to their full length (Sl analysis) if they extend into this downstream region. One experiment showed a possible protected region that terminated 50-100 bp downstream of the galk translation stop. The existence of a transcription terminator can be confirmed by conventional techniques by using a terminator probe system. The gal operon transcript clearly does not extend to the PvuII site (see, Table A, map position 8) because no full length protection of 5' labelled PvuII fragments occurs from that site.

fragments, fragment I, (map positions 4-6, See, Table A), and fragment II, (map positions 6-8, See Table A), and the insert of pSaulO were used as sources of probes for Sl walking from the 3' to 5' end of the message. All fragments through this region are protected, except the fragment containing the P2 promoter which shows partial and full protection. The complete protection from Sl digest indicates a polycistronic message which initiates upstream at Pl and continues to approximately 100 bp downstream of galK.

The above data is indirect evidence of a polycistronic mRNA of the <u>Streptomyces gal</u> operon. Sl analysis using a long contiguous DNA fragment (e.g., the 4.5 kb <u>HindIII-SacI</u> fragment, see map position 7 of Table A) has been used to confirm the transcript size.

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EXAMPLE 4

LOCALIZATION OF S. LIVIDANS GAL OPERON GALE AND GALT GENES

(i) Summary

The <u>S. lividans gal</u> operon <u>gal</u>E gene was localized to 1.5 Kb <u>PvuII</u> fragment (map position, 4-6 of Table A) of pLIVGAL1 (Figure 1).

The <u>S</u>. <u>lividans gal</u> operon <u>gal</u>E coding sequences extend through the <u>MluI</u> site (map position 5 of Table A).

The <u>S. lividans gal</u> operon <u>gal</u>T gene was localized within the 1.15 Kb <u>Nru-Pvu</u>II region (see, Table A, map positions la-4) of pSLIVGAL1.

The direction of \underline{S} . <u>lividans gal</u> operon <u>galE</u> and <u>galT</u> transcription is the same as \underline{galK} gene.

(ii) Experimental

It was necessary to identify the other functions contained on pLIVGAL1; specifically, does this plasmid encode for the enzyme galactose epimerase (galE) or the enzyme galactose transferase (galT). The Streptomyces gal operon galK gene was identified by its ability to complement an E. coli galK host. Thus, identification of the Streptomyces galT and galE genes was tested for by complementation of E. coli galE or galT hosts,

respectively. An <u>E. coli galT</u> strain (CGSC strain #4467, W3101) and two <u>galE</u> strains (CGSC strain #4473; W3109 and CGSC strain #4498; PL-2) were obtained to test for complementation by the pSLIVGAL1 clone.

The ca. 9 Kb <u>HindIII-SphI</u> fragment (see, Table A, map positions 1-16) containing the <u>Streptomyces lividans</u>

gal operon galk gene was inserted into pUC19. This fragment was situated within pUC19 such that transcription from the <u>Plac</u> promoter of pUC19 is in the same direction as the <u>Streptomyces galk gene</u>. pUC19 is described in Yanisch-Perrou, et al., <u>Gene</u>, <u>33</u>, 103 (1983).

Complementation was assayed by growth on MacConkeygalactose plates. Cells which can utilize galactose

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[galE+, galT+, galK+] will be red to pink on this medium. E. coli strain PL-2 (see, Example 2) containing pUC19 with the HindIII-SphI insert were pink on the indicator plate indicating that the HindIII-SphI fragment contains the Streptomyces lividans galE gene. gene was later mapped to within the 4.5 Kb HindIII-SacI (the SacI site is near the region around map position 7-8 of Table A) fragment. If the sequences from the MluI site (map position 5 of Table A) to the SacI site were removed 10 galE complementation of E. coli PL-2 was not detected. The 5' end of the galk gene is 70 base pairs (bp) from the MluI site. Therefore it seemed likely that the MluI site was contained within the 5' or 3' end of the galE gene. To determine the direction of galE transcription, the 15 HindIII-SacI fragment was inserted into pUC18. configuration, the Streptomyces lividans galk gene is in the opposite orientation with respect to Plac. HindIII-SphI clone did not complement E. coli PL-2 indicating the galE is transcribed in the same direction as galk. In addition it was concluded that the MluI site is contained within the 3' end of the galE gene. sequence analysis of the PvuII-MluI fragment (See, Table A, map position 4-5) has identified an open reading frame which encodes for a polypeptide of predicted molecular weight of 33,000 daltons. The 5' end of this reading frame is located approximately 176 bp from the PvuII site (See, Table A, map position 4). Therefore, the sequencing results support the conclusion that the 3' end of galE traverses the MluI site (see, Table A, map position 5).

Similar experiments to localize the galT gene on pSLIVGAL1 were attempted with the galT hosts.

The region between Pl and the 5' end of galE was sequenced to identify the galT gene. Translation of the DNA sequence to the amino acid sequence identified a reading frame which encodes a protein showing a region of homology to the yeast transferase.

EXAMPLE 5

GALACTOSE INDUCTION OF S. LIVIDANS GAL OPERON GALK GENE

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(i) Summary

Galactokinase expression is induced within one hour after the addition of galactose to culture medium.

Galactokinase expression is 10 times higher in the presence of galactose versus glucose or no additional carbon source within 6 hours after addition of the sugar.

(ii) Experimental

Galactose induction of the Streptomyces lividans galK gene was examined by assaying for galactokinase activity at 1, 3, 6 and 24 hours after the addition of Two liters of YM + 0.1M MOPS (pH 7.2) were inoculated with 2x10⁷ spores of <u>Streptomyces</u> <u>lividans</u> 1326. After 21 hours growth, galactose or glucose were added to a final concentration of 1%. One, three, six and twenty four hours after the addition of sugar, cells were isolated and assayed for galactokinase activity. Total RNA was prepared by procedures described in Hopwood et al., cited above.

An increase in galactokinase synthesis was observed one hour after the addition of galactose. increase continued over time (1 to 24 hours). Sl analysis of RNA isolated from the induced cultures confirmed that the increase in galk activity was due to increased levels of the Pl promoter transcript.

The Sl data and the induction studies suggest the 30 following model for gene expression within the Streptomyces gal operon. The Pl promoter is the galactose inducible promoter. The Pl transcript includes galT, galE and galK. The P2 promoter is constitutive and its transcript includes galE and galK.

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It is interesting to note that the E. coli gal operon also has two promoters, Pl and P2. [See, Nusso et al., Cell, 12, 847 (1977)]. Pl is activated by cAMP-CRP binding whereas P2 is inhibited by cAMP-CRP. Translation of the E. coli gal operon galE coding sequence is more efficient when transcription initiates at P2 which serves to supply a constant source of epimerase even in the absence of galactose or the presence of glucose [See, Queen et al., <u>Cell</u>, <u>25</u>, 241 (1981)]. The epimerase functions to convert galactose to glucose 1-phosphate during galactose utilization and convert UDP-glucose to UDP-galactose which is required for E. coli cell wall biosynthesis. It is possible that the P2 promoter of the Streptomyces galk operon also serves to supply epimerase and galactokinase in the absence of galactose or during secondary metabolism.

EXAMPLE 6

THE S. COELICOLOR GAL OPERON

(i) Summary

The restriction map of a fragment containing the \underline{S} . coelicolor galk gene is identical to the restriction map of the \underline{S} . lividans gal operon. (See, Figure 3).

S. coelicolor can grow on minimal media containing galactose as the sole carbon source.

Galactokinase expression in \underline{S} . coelicolor is induced by the addition of galactose to the growth media.

A promoter analogous and most likely identical to Pl is responsible for galactose induction of the \underline{S} . coelicolor gal operon.

(ii) Experimental

An approximately 14 kb partial <u>Sau</u>3A fragment containing the <u>S. coelicolor galK</u> gene was isolated by K. Kendall and J. Cullum at the University of Manchester Institute of Science and Technology, Manchester, UK

- l (unpublished data; personal communication). They were able to localize the S. coelicolor galk gene within a 3 kb ECORI fragment by complementation of a S. coelicolor galk mutant. The position of a number of restriction sites
- within the <u>S. lividans gal</u> operon are identical to those found within, upstream and downstream of the <u>EcoRI</u> fragment containing the <u>S. coelicolor galk</u> gene (Figure 3). Thus, it seems likely that the gene organization of the <u>S. coelicolor gal</u> operon is identical to the <u>S. lividans gal</u> operon.

Galactose induction of the S. coelicolor galk gene was examined by immunoblotting. S. coelicolor was grown in YM + 1% galactose or 1% glucose (Ymglu or Ymgal) for 20 hours at 28 C. Galactokinase expression was

15 detected using rabbit antisera prepared against purified

E. coli galactokinase. The protein detected was the approximate site of the E. coli and S. lividans galk gene product. Galactokinase expression is galactose induced since it was detected only when S. coelicolor was grown in 20 ym + galactose (Ymgal).

S1 nuclease protection studies were performed to determine if galactose induction of the <u>S. coelicolor gal</u> operon is directed by a promoter analogous to the <u>S. lividans</u> Pl promoter. RNA was isolated from S. coelicolor

- 25 grown in Ym + 1% galactose or 1% glucose (Ymgal or Ymglu). The hybridization probe used for Sl analysis of this RNA was a 410 bp Sau 3A fragment which contains the S. lividans Pl promoter, its transcription start site and the 5' end of the galT gene. The Sl protected fragment
- detected by this analysis co-migrated with the protected fragment detected when the probe was hybridized to RNA isolated from S. <u>lividans</u> grown in the presence of galactose. Thus, this result shows that galactose induction of the S. <u>coelicolor gal</u> operon is directed by a
- 35 sequence indistinguishable from the <u>S. lividans</u> Pl promoter.

It should be noted that the following strains of Streptomyces have been observed to be able to grow on medium containing galactose as the only carbon source:

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- S. <u>albus</u> J1074 (obtained from Dr. Chater, John Innes Foundation, Norwich, England)
- S. carzinostaticus ATCC accession number 15944
- S. carzinostaticus ATCC accession number 15945
- 10 S. antifibrinolyticus ATCC accession number 21869
 - S. antifibrinolyticus ATCC accession number 21870
 - S. antifibrinolyticus ATCC accession number 21871
 - S. longisporus ATCC accession number 23931

scope of the following claims.

The abbreviation "ATCC" stands for the American Type Culture Collection, Rockville, Maryland, U.S.A.

While the above descriptions and Examples fully describe the invention and the preferred embodiments thereof, it is understood that the invention is not limited to the particular disclosed embodiments. Thus, the invention includes all embodiments coming within the

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Claims for the Contracting States: BE, CH, DE, FR, GB, IT, LI, LU, NL, SE

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- 1. A recombinant DNA molecule comprising a Streptomyces gal operon or any regulatable and functional derivative thereof.
- 2. The molecule of Claim 1 wherein the operon is a S. lividans, S. coelicolor, S. azuraeus, S. albus, S. carzinostaticus, S. antifibrinolyticus or S. longisporus gal operon.
 - 3. The molecule of Claim 2 wherein the operon is a <u>S</u>. <u>lividans gal</u> operon.
 - 4. The molecule of Claim 3 which has the following coding sequence:

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-100 -120 -110 - 90 -80 -70 CTA CGC CTC CGC GTT CAG TAA TTG AAC ACT TTT GGT GAT GAA CTT TGT TTG ATT CTG 10 -60 -50 -40 -30 -20 ATC TCA CAC CCC CCT CCT CCC TCG TTG TCA TCT CTT ATC TTT GAT TCT CTT CCA TCA TTC -10 1 10 20 30 40 15 ACG GCC GTC CTG GTG ACT CAT GGG TGG GTG CAG AGG AGT GCG GCA GTG AAG AAG ACC Wet Thr His Gly Trp Val Gln Arg Ser Ala Ala Val Lys Lys Thr galT 70 50 60 80 90 100 TCG ACC CGG CTG GCC GAC GGC CGT GAG CTG GTC TAC TAC GAC CTG CGC GAC GAC ACC Ser Thr Arg Leu Ala Asp Gly Arg Clu Leu Val Tyr Tyr Asp Leu Arg Asp Asp Thr 20 110 120 130 140 150 GTG CGC GAC GCC GTG GAC CGC CGT CCG CTG GAG CGG ACC GTC ACC ACG TCC GAG GTG Val Arg Asp Ala Val Asp Arg Arg Pro Leu Glu Arg Thr Val Thr Thr Ser Glu Val 170 160 180 190 200 210 25 CGA CGC GAC CCG CTG CTC GGC GAC TCC GCG CCG TCG CGC CTC GCA CCG GCA GGC GCG Arg Arg Asp Pro Leu Leu Gly Asp Ser Ala Pro Ser Arg Leu Ala Pro Ala Gly Ala 220 230 240 250 260 CAC CTA CCA TCC GCC GGC CGA CCA GTG CCC GCT GTG CCc GTC GGA CGG GGA ACG GCT His Leu Pro Ser Ala Cly Arg Pro Val Pro Ala Val Pro Val Cly Arg Cly Thr Ala 30 280 290 300 310 320 330 GAG CGA GAT CCG GCC TAT GAC GTG GTG GTC TTC GAG AAT CGC TTT CCC TCG CTG GCC Glu Arg Asp Pro Ala Tyr Asp Val Val Val Phe Glu Asn Arg Phe Pro Ser Leu Ala

			3	40			350			360			3	70			380		
5	GGT Gly	GAC Asp	TCC Ser	GCG Gly	CGC Arg	TGC Cys	GAG Glu	GTC Val	GTC Val	TGC Cys	TTC Phe	ACC Thr	TCC Ser	GAC Asp	CAC His	GAC Asp	CCC Ala	TCC Ser	TTC Phe
	390)		4	00			410			420)		4	30			440	
	GCC Ala	GAC Asp	CTG Leu	AGC Ser	GAG Glu	GAG Glu	CAG Gln	GCC Ala	CGG Arg	CTG Leu	GTC Val	GTC Val	GAC Asp	GCC Ala	TGG Trp	ACG Thr	GAC Asp	CGC Arg	ACC Thr
10		450	o		4	60			470			480)		4	190			500
10	TCC Ser	GAG Glu	CTG Leu	TCC Ser	CAT His	CTG Leu	CCC Pro	TCC Ser	CTT Val	GAA Glu	CAG Gln	GTG Val	TTC Phe	TGC Cys	TTC Phe	GAG Glu	AAC Asn	CGG Arb	GCC Gly
			519	0		ţ	520			530			540)		5	550		
15	GCC Ala	GAG Glu	ATC Ile	CCC	CTG Val	ACG Thr	CTG Leu	GGT Gly	CAC <u>His</u>	CCC Pro	CAC His	GGG Gly	CAG Gln	ATC Ile	TAC Tyr	GCC Ala	TAC Tyr	CCG Pro	TTC Phe
	560)		57	0		į	580			590			60	0		1	510	
	ACC Thr	ACC Thr	CCC Pro	CGC Arg	ACC Thr	GCC Ala	CTG Leu	ATG Net	CTC Leu	CGT Arg	TCA Ser	CTC Leu	GCC	GCC Ala	CAC His	AAC Lys	GAC Asp	GCG Ala	ACG Thr
20		620)		63	0			640			650			66	0			670
	GG(G1 ₃	GGC Gly	G GGG	. AAC	CTC Leu	TTC Phe	GAC Asp	TCC Ser	GTG Val	CTG Leu	GAC Glu	GAG Glu	GAG Glu	CTG Leu	GCC	GGT Gly	GAG Glu	CGG Arg	GTC Val
			680)		69	0			700	;		710			72	0		
25	GT(Va	C CTC	G GAC	GCT Gly	GAC	CAC His	TGG Trp	GCC	GCC	TTC Phe	GTC Val	GCC Ala	TAC	GGC Gly	GCG	CAC	TCC Trp	CCG Pro	TAC Tyr
	730)		75	ю			760			770	i		78	0	
	GA Gl	G GT	G CAC	CTC Leu	TAC Tyr	C CCC	Lys	CGC Arg	CGG Arg	GTC Yal	Pro	GA1	CTG Leu	Lev	GGG	CTC Leu	GAC Asp	GAG Glu	GCG
30		790			800)		81	0			820			830)		84	0
30	GC Al	T CG	C AC	A GAJ	TTO Pho	C CCC	Lys	GT(TAC L Tyr	CTC Lev	GAC Glu	CTC	CTC Lev	AGC Arg	CGT	TTC Phe	GAC Asp	CGC Arg	ATC Ile

TTC GGC GAG GGC GAG CCC CCG ACC CCC TAC ATC GCG GCC TGG CAC CAG GCG CCC TTC Phe Gly Glu Gly Glu Pro Pro Thr Pro Tyr Ile Ala Ala Trp His Gln Ala Pro Phe GGG CAG CTG GAG TTC GAG GGT GTG ACG CGC GAC GAC TTC GCG CTC CAC CTG GAA CTT Gly Gln Leu Glu Phe Glu Gly Val Thr Arg Asp Asp Phe Ala Leu His Leu Glu Leu TTC ACT TCC GCC GTA CCT CCG GCA AGC TGA AGT TCC TCG CGG GCT CCG AAT CCG GCA Phe Thr Ser Ala Val Arg Pro Ala Ser --galP2 TGAACG TGTTCATCAA CGACGTACCC CCGGAGCGCC CGGCCGAGCG ACTGCGAGAG GTAGCGAG TTC ATG AGC GGG AAG TAC CTG GTG ACA GGT GGT GCC GGA TAC GTC GGC AGC GTC GTC Met Ser Gly Lys Tyr Leu Val Thr Gly Gly Ala Gly Tyr Val Gly Ser Val Val galE GCC CAG CAC TTG GTG GAG GCG GGG AAC GAG GTC GTG GTG CTG CAC AAT CTG TCG ACC Ala Cln His Leu Val Glu Ala Cly Asn Glu Val Val Leu His Asn Leu Ser Thr GGC TTC CGT GAG GTG TGC CGG CGG GTG CCT CGT TCG TCG AGG CGA CAT CCG GGA CGC Gly Phe Arg Clu Val Cys Arg Arg Val Pro Arg Ser Ser Arg Arg His Pro Gly Arg CGC CAA GTG CGT GGA CGG CTC TCG TTC GAC GGC GTG CTG CAC TTC GCC GCC TTC TCC Arg Gln Val Arg Gly Arg Leu Ser Phe Asp Gly Val Leu His Phe Ala Ala Phe Ser CAG GTC GGC GAG TCG GTC GTG AAG CCC GAG AAG TAC TGG GAC AAC AAC GTC GGT GGC Gln Val Gly Glu Ser Val Val Lys Pro Glu Lys Tyr Trp Asp Asn Asn Val Gly Gly

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	1370 ACC ATG GCG CTG				1	380			1390			14	00		1	410			1420
5	ACC Thr	ATG Met	GCG Ala	CTG Leu	CTG Leu	GAG Glu	GCC Ala	ATG Met	CGC	CCC Gly	GCG Ala	GGT Gly	GTG Val	CGG Arg	CGC Arg	CTC Leu	GTC Val	TTC Phe	TCC Ser
			14	130		1	440			1450)		14	60		1	470		
	TCC Ser	ACG Thr	GCC Ala	GCC Ala	ACG Thr	TAC Tyr	GGC Gly	GAG Glu	CCC Pro	GAG Glu	CAG Gln	GTT Val	CCC Pro	ATC Ile	GTC Val	GAG Glu	TCC Ser	GCG Ala	CCG Pro
10	1480)		14	190		1	500			1510)		15	520		1	530	
	ACG Thr	AGG Arg	CCC Pro	ACC Thr	AAT Asn	CCG Pro	TAC Tyr	GGC Gly	GCC Ala	TCG Ser	AAG Lys	CTC Leu	GCC Ala	GTC Val	GAC Asp	CAC His	ATG Net	ATC Ile	ACC Thr
		1540)		13	550		1	560			1570)		13	580		:	1590
15	GGC Gly	GAG Glu	GCG Ala	GCG Ala	GCC Ala	CAC His	GGG Gly	CTG Leu	GGC Gly	GCG Ala	GTC Val	TCC Ser	GTC Val	CCG Pro	TAC Tyr	TTC Phe	AAC Asn	GTC Val	GCG Ala
	1600			0		1	610			620			1630	0		1	640		
	GGC Gly	GCG Ala	TAC Tyr	GGG Gly	GAG Glu	TAC Tyr	GGC Gly	GAG Glu	CGC Arg	CAC His	GAC Asp	CCC Pro	GAG Glu	TCG Ser	CAT His	CTG Leu	ATT Ile	CCC Pro	CTC Leu
20	1650			166	0		1	670			1680			169	0		1	700	
	GTC Val	CTT Leu	CAA Gln	GTG Val	GCG Ala	CAG Gln	GGC Gly	AGG Arg	CGG Arg	GAG Glu	GCC Ala	ATC Ile	TCC Ser	GTC Val	TAC Tyr	GGC Gly	GAC Asp	GAC Asp	TAC Tyr
		1710	i		172	o*		1	730		;	1740			175	0		1	760
25	CCG Pro	ACC Thr	CCG Pro	GAC Asp	CGA	CCT Pro	GTC Val	TGC Cys	GCG	ACT Thr	ACA Thr	TCC Ser	ACG Thr	TCC Ser	CCG Pro	ACC Thr	TGG Trp	CCG Pro	AGC Arg
	1770					178				790			1800			181			
	CCC Pro	ACC Thr	TGC Cys	TGG Trp	CCG Pro	TCC Cys	Y)s	GCC	GCC	CCG Pro	GGC Gly	GAG Glu	CAC	CTC Leu	ATC	TGC	AAC	CTG Leu	GGC Gly
30	1820			1830			184			_	850			1860			187		
	AAC Asr	GGC Gly	AAC	GGC	Phe	TCC Ser	GTC Val	CGC Arg	GAG Glu	GTC Val	GTC Val	GAG Glu	ACC	GTG Val	CGG Arg	CGG	GTG Val	ACG Thr	GGC Gly

	18	880			1890			190	0		1	910			1920	l		193	0
5	CAT His	CCC Pro	ATC Ile	CCC Pro	GAG Glu	ATC Ile	ATG Met	GCC Ala	CCC Pro	CGC Arg	CGC Arg	GCC	CGC Arg	GAC Asp	CCC Pro	GCG	GTC Val	CTC Leu	CTC Val
		11	940			1950			196	0		1	970			1880			1990
	CCG Ala	TCC Ser	GCC Ala	GGC Gly	ACC Thr	GCC Ala	CCC Arg	GAG Glu	AAC Lys	CTG Leu	GGC Gly	TGG Trp	AAC Asn	CCC Pro	TCC Ser	CGC Arg	GCG Ala	GAC Asp	CTC Leu
10			2	000		;	2010			2020	0		2	030			2040		
	GCC Ala	ATC Ile	GTG Val	TCG Ser	GAC Asp	GCG Ala	TGG Trp	GAC Glu	TTG Leu	CCC Pro	CAG Gln	CGG Arg	CGC Arg	GCG Ala	GGC Gly	CAC Gln	TAG	ΤA	
		20	050		,	2060			207	0		2	080			2090			2100
15	ACC	GCA	CTT	ACC	GGA	AAG	GCG	ACC	CGT	CAG	GGC	Met	Gly	GAG Glu	GCT Ala	GTC Val	GGG Gly	GAA Glu	CCG Pro
		2110				:	2120			2130)	gal		140		:	2150		
	TCC	GCG	AGC	GGT	TCC	GGG	AGC	TCT	ACG	GGG	CGG	AGC	CCC	AGG	GGG	TGT	GGG	CGC	CGA
20	2160		Ser		3er 170	GIY		2180	inr	GIY			Arg			Cys		_	Arg
	•				-			•			2190				200			2210	
	GCG Ala	GCC Gly	CGG Arg	GAG Glu	AAC Asn	CTC Leu	ATC Ile	GGG Gly	GAG <u>Clu</u>	CAC His	ACC Thr	GAC Asp	TAC Tyr	AAC Asn	CAC Asp	GGC Gly	TTC Phe	GTC Val	ATG Met
		2220)		23	230		:	2240	;	ļ	2250)		23	260		2	270
25	CCT Pro	TCG Ser	CCC Pro	TGC Cys	CGC Arg	ACC Thr	AGC Arg	TCG Ser	CGG Arg	CCG Pro	TCT Ser	CCC Pro	GGC Gly	GCG Ala	AAC Asn	GAC Asp	GGC Gly	ATC Ile	CTG Leu
			2280)		22	90	•	2	300			2310)		23	20		
30	CGC Arg	CTG Leu	CAC	TCC Ser	GCC Ala	GAC Asp	GTC Val	GAC Asp	GCC Ala	GAC Asp	CCG Pro	GTC Val	GAG Glu	CTG Leu	CGC Arg	GTC Val	GCC Ala	GAC Asp	CTG Leu
	2330			2340)		23	50		2	360			2370	•		23	80	
	GCC Ala	CCC Pro	GCG Ala	TCG Ser	GAC Asp	AAG Lys	TCC Ser	TGC Trp	ACG Thr	GCC Ala	TAC Tyr	CCC Pro	TCG Ser	GGC Gly	GTC Val	CTG Leu	TGG Trp	GCG Ala	CTG Leu

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	23	80			2400			24	0		2	420			2430			24	40
5	CGC G	AG G	CC (GGA Gly	CAC His	GAG Glu	CTG Leu	ACC (GGC Gly	GCC Ala	GAC Asp	GTC Val	CAC	CTG Leu	GCC Ala	TCC Ser	ACC Thr	GTC Val	CCG Pro
		24	150			2460	}		24	70		2	480			2490)		
	TCC G	GG (CCC	CCC Gly	CTC Leu	TCC Ser	TCC Ser	TCC Ser	GCG Ala	GCC Ala	CTG Leu	GAG Glu	CTC Val	CCT Arg	CCC Pro	CTG Leu	GCG Ala	ATG Wet	AAC Asn
10	2500			510			2520				530			540			2550		
•	GAC (CTG '	TAC Tyr	GCC Ala	CTC Leu	GCG Ala	CTG Leu	CGC Arg	GGC Gly	TGG Trp	CAG Gln	CTG Leu	GCC Ala	CCC Arg	CTG Leu	TCC Cys	CAG Gln	CGC Arg	CCC Ala
	250		•		2570			2580				590			2600			261	
15	GAG Glu	AAC Asn	GTC Yal	TAC Tyr	GTC Val	GGC Gly	GCC Ala	CCC Pro	GTC Val	GGC Gly	ATC Ile	ATG Met	GAC Asp	CAG Gln	ACG Thr	GCG Ala	TCC Ser	GCC	TGC Cys
		26	20			2630			2640)		2	650			2660			2670
20	TGC Cys	GAG Glu	GCC Ala	GGC Gly	ACG Thr	CCC Pro	TCT Ser	TCC Ser	TCG Ser	ACA Thr	CCC	GCG	ACC Thr	TCT Ser	CCC Pro	ACC Ser	GGC Gly	ACA Arg	TCC Ser
20			2	680			2690			270	Ю		2	710			2720		
	CCT Pro	TCG Ser	ACC Thr	TCG Ser	CCG Pro	CCC Pro	AGG Arg	GGA Gly	TGC Cys	GCC	TGC Cys	TGC Trp	TCG Ser	TCG Ser	ACA	CCC Pro	GGG	TCA Ser	AGC Ser
	2730	0		2	740			2750			,276	O		2	770			2780	•
25	ACT Thr	CCC Pro	ACA Thr	GC0	AGC Are	G GCC	AGT Ser	ACG Thr	GCA Ala	AGC Set	GCC - Alz	GCC	CGG Arg	GCT	GCC	AGA L Are	AGG Arg	GCC Ala	CCG Pro
		279	0		:	2800			2810)		282	20		2	2830			2840
	Arg	TGC Cys	TGC	GCC Al:	TC0	G ACC	G CGC	TGC Cys	GAC Asp	CT(CCC l Pro	TAC Ty:	G GCC	. Ast	CTC Lei	G GAC	GCC Ala	GCC Ala	CTG Leu
30			283				2860			287			288				2890		
30	GAG Glu	CGC Arg	CTO Let	G GG G G1	C GA	C GA	G GAG	G GAC	GTC	G CG	C CG	C CT	G GT(C CGC	G CAG	C GT(GTC I Val	ACC Thi	GAG Glu

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	2900 2910						29	20		:	2930			2940)		29	50	
5	GAC (
	29	960			2970	į		29	980		:	2990			3000)		30	010
• •	GCG G Ala F																		
10		3	1020			3030)		30	140		:	3050			3060)		
	CGA (
	3070		;	3080			3090)		3	100		;	3110			3120)	
15	ACC (
	313	30		;	3140			3150)		3	60		:	3170			3180)
20	Ala '																		
20		31	90		;	3200			3210)		32	220		:	3230			3240
	GTG T																		
			32	250		;	3260			3270)		32	280		:	3290		
25	GCT :								TCC	ccc	GCG	GGT	AGT	CGG	GGA	TCA	ccc	ACA	TGA
	3300																		
	GCT	CCT	AGC	CGC				•											

- 5. The molecule of Claim 1 which further comprises a foreign functional DNA sequence operatively linked to such operon.
 - 6. A transformed host microorganism or cell comprising the molecule of Claim 5.
 - 7. A method of preparing a transformed host microorganism or cell comprising the molecule of Claim 5 which comprises transforming an appropriate host microorganism or cell with such molecule.
- 8. A recombinant DNA vector comprising the molecule of Claim 5, and, optionally, additionally comprising a replicon.
 - 9. A transformed host microorganism or cell comprising the recombinant DNA vector of Claim 8.
- 10. A method of preparing a transformed host microorganism or cell comprising the recombinant DNA vector of Claim 8 which comprises transforming an appropriate host microorganism or cell with such vector.
- 20 DNA sequence which comprises cultivating a transformed host microorganism or cell comprising the recombinant DNA vector of Claim 8 under suitable conditions such that the functional DNA sequence is expressed.
- 12. A method of regulating the expression of a

 25 foreign functional DNA sequence which comprises
 cultivating a transformed host microorganism or cell which
 contains the recombinant DNA vector of Claim 8 under
 appropriate conditions such that expression of the
 sequence is regulatable.
- 30 . 13. A recombinant DNA molecule comprising a Streptomyces gal operon P2 promoter expression unit or any functional derivative thereof.
 - 14. The molecule of Claim 13 wherein the expression unit is a \underline{S} . lividans, \underline{S} . coelicolor, \underline{S} .
- 35 <u>azuraeus</u>, <u>S</u>. <u>albus</u>, <u>S</u>. <u>carzinostaticus</u>, <u>S</u>. <u>antifibrinolyticus</u> or <u>S</u>. <u>longisporus gal</u> operon P2 promoter expression unit.

i... One molecule of Claim 14 with is a S. lividans gal operon P2 promoter expression unit.

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- 16. The molecule of Claim 13 which further comprises a foreign functional DNA sequence operatively linked to such expression unit.
- 17. A transformed host microorganism or cell comprising a recombinant DNA molecule wherein such molecule comprises the molecule of Claim 16.
- 18. A method of preparing a transformed host
 microorganism comprising the molecule of Claim 16 which
 comprises transforming an appropriate host microorganism
 or cell with such molecule.
 - 19. A recombinant DNA vector comprising the molecule of Claim 16, and, optionally, additionally comprising a replicon.
 - 20. A transformed host microorganism or cell comprising the recombinant DNA vector of Claim 19.
 - 21. A method of preparing a transformed host microorganism or cell comprising the recombinant DNA vector of Claim 19 which comprises transforming an appropriate host microorganism with such vector.
 - DNA sequence which comprises cultivating a transformed host microorganism or cell comprising the recombinant DNA vector of Claim 19 under suitable conditions such that the functional DNA sequence is expressed.
 - 23. A recombinant DNA molecule comprising a Streptomyces gal operon Pl promoter regulated region or any regulatable and functional derivative thereof.
 - 24. The molecule of Claim 23 wherein the region is a S. lividans, S. coelicolor, S. azuraeus, S. albus, S. carzinostaticus, S. antifibrinolyticus or S. longisporus gal operon Pl promoter regulated region.
- 25. The molecule of Claim 24 wherein the region is a S. <u>lividans gal</u> operon Pl promoter regulated region.

- 26. The molecule of Claim 23 which further comprises a foreign functional DNA sequence operatively linked to such regulated region.
- 27. A transformed host microorganism or cell comprising the molecule of Claim 26.
 - 28. A method of preparing a transformed host microorganism or cell comprising the molecule of Claim 26 which comprises transforming an appropriate host microoganism or cell with such molecule.
- 29. A recombinant DNA vector comprising the molecule Claim 26, and, optionally, additionally comprising a replicon.
 - 30. A transformed host microorganism or cell comprising a recombinant DNA vector of Claim 29.
- 15 31. A method of preparing a transformed host microorganism or cell comprising the recombinant DNA vector of Claim 29 which comprises transforming an appropriate host microorganism or cell with such vector.
- 20 DNA sequence which comprises cultivating a transformed host microorganism or cell comprising the recombinant DNA vector of Claim 29 under suitable conditions such that the functional DNA sequence is expressed.
- 33. A method of regulating the expression of a foreign functional DNA sequence which comprises cultivating a transformed host microorganism or cell which contains the recombinant DNA vector of Claim 29 under appropriate conditions such that expression of the sequence is regulatable.
- 34. A recombinant DNA molecule comprising a Streptomyces gal operon P2 promoter or any functional derivative thereof.
 - 35. The molecule of Claim 34 wherein the promoter is a <u>S</u>. <u>lividans</u>, <u>S</u>. <u>coelicolor</u>, <u>S</u>. <u>azuraeus</u>, <u>S</u>. <u>albus</u>, <u>S</u>. <u>carzinostaticus</u>, <u>S</u>. <u>antifibrinolyticus</u> or <u>S</u>. <u>longisporus</u>

gal operon P2 promoter.

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- promoter is a S. <u>lividans gal</u> operon P2 promoter.
- 37. The molecule of Claim 34 which further comprises a foreign functional DNA sequence operatively linked to the P2 promoter.
 - 38. A transformed host microorganism or cell comprising the molecule of Claim 37.
- 39. A method of preparing a transformed host microorganism or cell comprising the molecule of Claim 37 which comprises transforming an appropriate host microorganism or cell with such molecule.
- 40. A recombinant DNA vector comprising the molecule of Claim 37 and, optionally, additionally comprising a replicon.
- 41. A transformed host microorganism or cell comprising the recombinant DNA vector of Claim 40.
 - 42. A method of preparing a transformed host microorganism or cell comprising the recombinant DNA vector of Claim 40 which comprises transforming an appropriate host microorganism with such vector.
 - 43. A method of expressing a foreign functional DNA sequence which comprises cultivating a transformed host microorganism or cell comprising the recombinant DNA vector of Claim 40 under suitable conditions such that the functional DNA sequence is expressed.
 - 44. A recombinant DNA molecule comprising a Streptomyces gal operon Pl promoter or any regulatable and functional derivative thereof.
- 45. The molecule of Claim 44 wherein the
 promoter is a S. <u>lividans</u>, S. <u>coelicolor</u>, S. <u>azuraeus</u>, S.
 <u>albus</u>, S. <u>carzinostaticus</u>, S. <u>antifibrinolyticus</u> or S.
 <u>longisporus gal</u> operon Pl promoter.
 - 46. The molecule of Claim 45 wherein the promoter is a \underline{S} . lividans gal operon Pl promoter.
- 35 47. The molecule of Claim 44 which further comprises a foreign functional DNA sequence operatively linked to the Pl promoter.

- 1 48. A transformed host microorganism or cell comprising the molecule of Claim 47.
 - 49. A method of preparing a transformed host microorganism or cell comprising molecule of Claim 47 which comprises transforming an appropriate host microorganism or cell with such molecule.
 - 50. A recombinant DNA vector comprising the molecule of Claim 47, and, optionally, additionally comprising a replicon.
- 51. A transformed host microorganism or cell comprising the recombinant DNA vector of Claim 50.
 - 52. A method of preparing a transformed host microorganism or cell comprising the recombinant DNA vector of claim 50 which comprises transforming an appropriate host microorganism with such vector.
 - 53. A method of expressing a foreign functional DNA sequence which comprises cultivating a transformed host microorganism or cell comprising the recombinant DNA vector of Claim 50 under suitable conditions such that the functional DNA sequence is expressed.
 - 54. A method of regulating the expression of a foreign functional DNA sequence which comprises cultivating a transformed host microorganism or cell which contains the recombinant DNA vector of Claim 50 under appropriate conditions such that expression of the sequence is regulatable.
 - 55. A recombinant DNA molecule comprising a Streptomyces gal operon galE gene, or any functional derivative thereof.
 - 30 56. The molecule of Claim 55 wherein the gene is a <u>S. lividans</u>, <u>S. coelicolor</u>, <u>S. azuraeus</u>, <u>S. albus</u>, <u>S. carzinostaticus</u>, <u>S. antifibrinolyticus</u> or <u>S. longisporus</u> gal operon galE gene.
 - 57. The molecule of Claim 56 wherein the gene is a <u>S</u>. <u>lividans gal</u> operon <u>gal</u>E gene.

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- 5d. The molecule of Claim 55 which further comprises a foreign functional DNA sequence operatively linked to the galE gene.
- 59. A transformed host microorganism or cell comprising the molecule of Claim 58.
- 60. A method of preparing a transformed host microoganism or cell comprising the molecule of Claim 58 which comprises transforming an appropriate host microorganism or cell with such molecule.
- 61. A recombinant DNA molecule comprising a <u>Streptomyces gal</u> operon <u>gal</u>T gene, or any functional derivative thereof.
 - 62. The molecule of Claim 61 wherein the gene is a S. lividans, S. coelicolor, S. azuraeus or S. albus, S. carzinostaticus, S. antifibrinolyticus and S. longisporus gal operon galT gene.
 - 63. The molecule of Claim 62 wherein the gene is a S. lividans gal operon galT gene.
- 20 comprises a foreign functional DNA sequence operatively linked to the <u>gal</u>T gene.
 - 65. A transformed host microorganism or cell comprising the molecule of Claim 64.
- 25 microorganism or cell comprising the molecule of Claim 64 which comprises transforming an appropriate host microorganism or cell with such molecule.
 - 67. A recombinant DNA molecule comprising a Streptomyces lividans gal operon galK gene, or any functional derivative thereof.
 - 68. The molecule of Claim 67 wherein the gene is a <u>S. lividans</u>, <u>S. coelicolor</u>, <u>S. azuraeus</u>, <u>S. albus</u>, <u>S. carzinostaticus</u>, <u>S. antifibrinolyticus</u> or <u>S. longisporus</u> gal operon galk gene.
- 35 69. The molecule of Claim 68 wherein is a \underline{S} . lividans gal operon galK gene.

- 70. The molecule of Claim 67 which further comprises a foreign functional DNA sequence operatively linked to the galk gene.
- 71. A transformed host microorganism or cell comprising the molecule of Claim 70.
 - 72. A method of preparing a transformed host microorganism or cell comprising the molecule of Claim 70 which comprises transforming an appropriate host microorganism or cell with such molecule.
- 10 73. A method of enabling a nongalactose utilizing host microorganism or cell to utilize galactose which comprises transforming such host with a recombinant DNA vector or molecule comprising a Streptomyces gal operon, or any portion of the Streptomyces gal operon
- which is adequate to enable such transformed host to utilize galactose, or any functional derivative thereof.
 - 74. A transformed host prepared by the method of Claim 73.

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Claims for the Contracting States: AT, ES, GR

 A method of preparing a transformed host microorganism or cell comprising the molecule which has the following sequence:

-70 -80 -90 -100 -110 -120 10 CTA CGC CTC CGC GTT CAG TAA TTG AAC ACT TTT GGT GAT GAA CTT TGT TTG ATT GTG -20 -30 -40 -50 -60 galPi 15 40 30 20 10 -10 ACC GGC GTC CTG GTG ACT CAT GGG TGG GTG CAG AGG AGT GCG GCA GTG AAG AAG ACC Met Thr His Gly Trp Val Gln Arg Ser Ala Ala Val Lys Lys Thr galT 90 100 80 70 60 50 TCG ACC CGG CTG GCC GAC GGC CGT GAG CTG GTC TAC TAC GAC CTG CGC GAC GAC ACC Ser Thr Arg Leu Ala Asp Gly Arg Clu Leu Val Tyr Tyr Asp Leu Arg Asp Asp Thr 20 140 150 130 120 110 GTG CGC GAC GCC GTG GAC CGC CGT CCG CTG GAG CGG ACC GTC ACC ACG TCC GAG GTG Val Arg Asp Ala Val Asp Arg Arg Pro Leu Glu Arg Thr Val Thr Thr Ser Glu Val 200 210 190 180 170 160 25 CGA CGC GAC CCG CTG CTC GGC GAC TCC GCG CCG TCG CGC CTC GCA CCG GCA GGG GCG Arg Arg Asp Pro Leu Leu Gly Asp Ser Ala Pro Ser Arg Leu Ala Pro Ala Gly Ala 260 250 240 230 220 CAC CTA CCA TCC GCC GGC CGA CCA GTG CCC GCT GTG CCC GTC GGA CGC GGA ACG GCT His Leu Pro Ser Ala Gly Arg Pro Val Pro Ala Val Pro Val Gly Arg Gly Thr Ala 330 320 30 310 300 290 280 GAG CGA GAT CCG GCC TAT GAC GTG GTG GTC TTC GAG AAT CGC TTT CCC TCG CTG GCC Glu Arg Asp Pro Ala Tyr Asp Val Val Val Phe Glu Asn Arg Phe Pro Ser Leu Ala

GGT GAC TOO GGG CGC TGC GAG GTC GTC TGC TTC ACC TCC GAC CAC GAC GCC TCC TTC Gly Asp Ser Gly Arg Cys Glu Val Val Cys Phe Thr Ser Asp His Asp Ala Ser Phe GCC GAC CTG AGC GAG GAG CAG GCC CGG CTG GTC GTC GAC GCC TGG ACG GAC CGC ACC Ala Asp Leu Ser Glu Glu Gln Ala Arg Leu Val Val Asp Ala Trp Thr Asp Arg Thr TCC GAG CTG TCC CAT CTG CCC TCC GTT GAA CAG GTG TTC TGC TTC GAG AAC CGG GGC Ser Glu Leu Ser His Leu Pro Ser Val Glu Gln Val Phe Cys Phe Glu Asn Arg Gly GCC GAG ATC GGG GTG ACG CTG GGT CAC CCG GAC GGG CAG ATC TAC GCC TAC CCG TTC Ala Clu Ile Cly Val Thr Leu Cly His Pro His Cly Cln Ile Tyr Ala Tyr Pro Phe ACC ACC CCC CGC ACC GCC CTG ATG CTC CGT TCA CTC GCC GCC CAC AAG GAC GCG ACG Thr Thr Pro Arg Thr Ala Leu Wet Leu Arg Ser Leu Ala Ala His Lys Asp Ala Thr GGC GGG GGG AAC CTG TTC GAC TCC GTG CTG GAG GAG GAG CTG GCC GGT GAG CGC GTC Gly Gly Gly Asn Leu Phe Asp Ser Val Leu Glu Glu Glu Leu Ala Gly Glu Arg Val GTC CTG GAG GGT GAG CAC TGG GCC GCC TTC GTC GCG TAC GGC GCG CAC TGG CCG TAC Val Leu Glu Gly Glu His Trp Ala Ala Phe Val Ala Tyr Gly Ala His Trp Pro Tyr GAG GTG CAC CTC TAC CCG AAG CGG CGG GTG CCC GAT CTG CTC GGG CTC GAC GAG GCG Glu Val His Leu Tyr Pro Lys Arg Arg Val Pro Asp Leu Leu Gly Leu Asp Glu Ala GCT CGC ACA GAA TTC CCC AAG GTC TAC CTG GAG CTG CTG AGG CGT TTC GAC CGG ATC Ala Arg Thr Glu Phe Pro Lys Val Tyr Leu Glu Leu Leu Arg Arg Phe Asp Arg Ile

TTC GGC GAG GGC GAG CCC CCG ACC CCC TAC ATC GCG GCC TGG CAC CAG GCG CCG TTC Phe Gly Glu Gly Glu Pro Pro Thr Pro Tyr Ile Ala Ala Trp His Gln Ala Pro Phe GGG CAG CTG GAG TTC GAG GGT GTG ACG CGC GAC GAC TTC GCG CTC CAC CTG GAA CTT Gly Gln Leu Glu Phe Glu Gly Val Thr Arg Asp Asp Phe Ala Leu His Leu Glu Leu TTC ACT TCC GCC GTA CGT CCG GCA AGC TGA AGT TCC TCG CGG GCT CCG AAT CCG GCA galP2 Phe Thr Ser Ala Val Arg Pro Ala Ser ---TGAACG TGTTCATCAA CGACGTACCC CCGGAGCGCG CGGCCGAGCG ACTGCGAGAG GTAGCGAG TTC ATG AGC GGG AAG TAC CTG GTG ACA GGT GGT GCC GGA TAC GTC GGC AGC GTC GTC Met Ser Gly Lys Tyr Leu Val Thr Gly Gly Ala Gly Tyr Val Gly Ser Val Val galE GCC CAG CAC TTG GTG GAG GCG GGG AAC GAG GTC GTG GTG CTG CAC AAT CTG TCG ACC Ala Cln His Leu Val Clu Ala Cly Asn Clu Val Val Val Leu His Asn Leu Ser Thr GGC TTC CGT GAG GTG TGC CGG CGG GTG CCT CGT TCG TCG AGG CGA CAT CCG GGA CGC Gly Phe Arg Glu Val Cys Arg Arg Val Pro Arg Ser Ser Arg Arg His Pro Gly Arg CGC CAA GTG CGT GGA CGG CTC TCG TTC GAC GGC GTG CTG CAC TTC GCC GCC TTC TCC Arg Gln Val Arg Gly Arg Leu Ser Phe Asp Gly Val Leu His Phe Ala Ala Phe Ser CAG GTC GGC GAG TCG GTC GTG AAG CCC GAG AAG TAC TGG GAC AAC AAC GTC GGT GGC Gln Val Gly Glu Ser Val Val Lys Pro Glu Lys Tyr Trp Asp Asn Asn Val Gly Gly

1

		1300	1390	1400	1410	1420
5	ACC ATG GCG CTG CTG Thr Wet Ala Leu Leu	GAG GCC ATG	CGC GGC GCG	GGT GTG CGG	CGG CTC GTC Arg Leu Val	TTC TCC Phe Ser
	1430	1440	1450	1460	1470	
	TCC ACG GCC GCC ACC Ser Thr Ala Ala Thi	TAC GGC GAG	CCC GAG CAG Pro Glu Gln	CTT CCC ATC Val Pro Ile	GTC GAG TCC Val Glu Ser	GCG CCG Ala Pro
10	1480 1490	1500		_		530
10	ACG AGG CCC ACC AA' Thr Arg Pro Thr As	T CCG TAC GGC n Pro Tyr Gly	GCC TCG AAG Ala Ser Lys	CTC GCC GTC Leu Ala Val	GAC CAC ATG Asp His Wet	ATC ACC Ile Thr
	1540	1550	1560	1570	1580	1590
15	GGC GAG GCG GCG GC Gly Glu Ala Ala Al	C CAC GGG CTG a His Gly Leu	GGC GCG GTC	C TCC GTG CCG Ser Val Pro	TAC TTC AAC Tyr Phe Asn	GTC GCG Val Ala
	1600	1610	1620	1630	1640	
	GGC GCG TAC GGG GA	G TAC GGC GAC	CGC CAC GAC	CCC GAG TCG Pro Glu Ser	CAT CTG ATT His Leu Ile	CCG CTG Pro Leu
20	1650 1660	1670	1680	_	-	700
	GTC CTT CAA GTG GG Val Leu Gln Val A	CG CAG GGC AGG	G CGG GAG GCG g Arg Glu Al	C ATC TCC GTC a lle Ser Val	TAC GGC GAC Tyr Gly Asp	GAC TAC Asp Tyr
	1.10	120	1730	1740	1750	1760
25	. CCG ACG CCG GAC C Pro Thr Pro Asp A	GA CCT GTG TG rg Pro Val Cy	C GCG ACT AC	A TCC ACC TCC	CCG ACC TGG	CCG AGG
	1770	1780	1790	1800	1810	
	CCC ACC TGC TGG C Pro Thr Cys Trp P	CCC TCC GCC GC Pro Cys Ala Al	CC GCC CCG GC la Ala Pro Gl	GC GAG CAC CTO Ly Glu His Le	4 11 2 2 3 3 3 3 3 3 3 3 3 3	•
30	1820 1830	1840	1850			
	AAC GGC AAC GGC T Asn Gly Asn Gly F	TTC TCC GTC CC The Ser Val A	GC GAG GTC GT rg Glu Val Va	TC GAG ACC GT al Glu Thr Va	G CGG CGG GT l Arg Arg Va	G ACG GGC

CAT CCG ATC CCC GAG ATC ATG GCC CCC CGC CGC GGC GGC GAC CCG GCG GTC CTG GTC His Pro Ile Pro Glu Ile Met Ala Pro Arg Arg Gly Arg Asp Pro Ala Val Leu Val GCG TCC GCC GGC ACC GCC GGC GAG AAC CTG GGC TGG AAC CCG TCC GGC GAC CTC Ala Ser Ala Gly Thr Ala Arg Glu Lys Leu Gly Trp Asn Pro Ser Arg Ala Asp Leu CCC ATC GTG TCG GAC GCG TGG GAG TTC CCG CAG CGG CGC GCC GGC CAG TAG TA Ala Ile Val Ser Asp Ala Trp Glu Leu Pro Gln Arg Arg Ala Gly Gln ---ACC GCA GTT ACC GGA AAG GCG AGG GGT CAG GGC ATG GGC GAG GCT GTC GGG GAA CCG Met Gly Glu Ala Val Gly Glu Pro TCG GCG AGC GGT TCC GGG AGC TGT ACG GGG CGG AGC CGG AGG GGG TGT GGG CGC CGA Ser Ala Ser Gly Ser Gly Ser Cys Thr Gly Arg Ser Arg Arg Gly Cys Gly Arg Arg GCG GGC CGG GAG AAC CTC ATC GGG GAG CAC ACC GAC TAC AAC GAC GGC TTC GTC ATG Ala Cly Arg Clu Asn Leu Ile Cly Clu His Thr Asp Tyr Asn Asp Cly Phe Val Met CCT TCG CCC TGC CGC ACC AGG TCG CGG CCG TCT CCC GGC GCG AAC GAC GGC ATC CTG Pro Ser Pro Cys Arg Thr Arg Ser Arg Pro Ser Pro Gly Ala Asn Asp Gly Ile Leu 2290 -CCC CTG CAC TCG GCC GAC GTC GAC GCC GAC CCG GTC GAG CTG CGC GTC GCC GAC CTG Arg Leu His Ser Ala Asp Val Asp Ala Asp Pro Val Glu Leu Arg Val Ala Asp Leu GCC CCC GCG TCG GAC AAG TCC TGG ACG GCG TAC CCC TCG GGC GTC CTG TGG GCG CTG Ala Pro Ala Ser Asp Lys Ser Trp Thr Ala Tyr Pro Ser Gly Val Leu Trp Ala Leu

	2	390			2400)		24	10		2	420			2430			24	40
5	CGC Arg	GAG Glu	GCC	GGA Gly	CAC His	GAG Glu	CTG Leu	ACC Thr	GGC Gly	GCC Ala	GAC Asp	CTC Val	CAC His	CTG Leu	GCC Ala	TCG Ser	ACC Thr	GTC Val	CCC Pro
			2450			2460				70			2480			2490			
	TCC Ser	GGG Gly	GCC Al:	GCC Gly	CTC Leu	TCC <u>Ser</u>	TCC Ser	TCC Ser	GCG Ala	GCC Ala	CTG Leu	GAG Glu	GTC Val	CGT Arg	CCC Pro	CTG Leu	GCG Ala	ATG Net	AAC Asn
10 2	500			2510			2520				30			540			2550		
	GAC Asp	CTC Lev	TA:	C GCC	CTC	GCG Ala	CTG Leu	CGC Arg	GGC Gly	TGG Trp	CAG Gln	CTG Leu	GCC Ala	CCC Arg	CTG Leu	TGC Cys	CAG Gln	CGC Arg	GCG Ala
2560			2570			2580				2590			2600				2610		
15		•	C GT	C TA	C GTC r Val	GGC Gly	GCC	CCC Pro	GTC Val	GGC Gly	ATC Ile	ATG Net	GAC Asp	CAG Gln	ACG Thr	GCG Ala	TCC Ser	CCC Ala	TGC Cys
	2620							2640 2650			2660				2670				
20	TGC Cys		•		C ACC	ccc r Pro	TCT Ser	TCC Ser	TCG Ser	ACA Thr	CCC	GCG	ACC Thr	TCT Ser	CCC	AGC Ser	GGC Gly	AGA Arg	TCC Ser
20				2680)		2690)		270	0		2	710			2720		
	CC1	T TC	G AC	CC TC	G CC	G CCC	AGG Arg	GGA Gly	TGC	GCC	TGC Cy≤	TGC Trp	C Ser	TCC Ser	ACA Thr	CCC Pro	GGG Gly	TCA Ser	AGC Ser
	273				2740			2750			1276	_			2770			2780	
25	AC Th	· T CC r Pi	CC A	CA G	CG AG	G GC	G AGT a Set	Thz	GCA - Ala	AGC Ser	GCC	C GCG	CGC a Arg	GCT Ala	GC(AG/	AGC Arg	GCC	CCG Pro
		27	790			2800			2810)		28	20			2830			2840
30	CG Ar	C TO	GC T	CG G	CC TC la Se	G AC	G CG	C TG(g Cy:	C GAC	GTC Val	CC Pr	G TA	C GCC	GA(C CTO	G GAG	C GCC	GCC Al:	CTG Leu
				850			2860			2870			288				2890		
	GA G1	.G C	GG C rg L	TG G	GC GA	C GA	G GA	G GA	G-GTO	C CGC	C CG g Ar	C CT g Le	G GTG	C CG	G CA g Hi	C GT s Va	C CTC	ACC Th	C GAG

GAC GAG CGC GTC GAA CGC GTC GTC GCG CTG GTG GAG TCG GCG ACA CCC GGC GCA TCG Asp Glu Arg Val Glu Arg Val Val Ala Leu Leu Glu Ser Ala Thr Pro Gly Ala Ser 297U GCG CCG TCC TGG TCC AGG GCC ACG CCT GCT GCG CGA CGA CTT CCG CAT CTC CTG CCC Ala Pro Ser Trp Ser Arg Ala Thr Pro Ala Ala Arg Arg Leu Pro His Leu Leu Pro Arg Ala Gly Pro Gly Arg Arg His Gly Pro Gly Leu Arg Gly Pro Arg Arg Met ACC GGC GGC GGC TTC GGC GGC TCG GCG ATC GTC CTG GTG GAG GCC GCC GCG GTG GAC 1.5 Thr Gly Gly Phe Gly Gly Ser Ala Ile Val Leu Val Glu Ala Ala Ala Val Asp GCC GTC ACC AAG GCC GTC GAG GAC GCC TTC GCC GCG GCC GCC CTC AAG CGT CCG CGC Ala Val Thr Lys Ala Val Glu Asp Ala Phe Ala Ala Ala Gly Leu Lys Arg Pro Arg GTG TTC GAG GCG GTG CCT CGC CCG GCC GCG CCT GGT CTG ACG GTC AGC CGA GCC Val Phe Glu Ala Val Pro Arg Arg Gly Ala Ala Pro Gly Leu Thr Val Ser Arg Ala GCT TCA CCA GCG TGT ACT CCG TGA TCC CCG GCG GGT AGT CGG GGA TCA CGC ACA TGA Ala Ser Pro Ala Cys Thr Pro ---GCT GCT AGC CGC which comprises transforming an appropriate host micro-

organism or cell with such molevule.

- 2. A method of preparing a transformed host microorganism or cell comprising a recombinant DNA vector comprising the sequence of Claim 1 which comprises transforming an appropriate host microorganism or cell 5 with such vector.
- 3. A method of expressing a foreign functional DNA sequence which comprises cultivating a transformed host microorganism or cell comprising the recombinant DNA vector of Claim 2 under suitable conditions such that the functional DNA sequence is expressed.
- 4. A method of regulating the expression of a foreign functional DNA sequence which comprises cultivating a transformed host microorganism or cell which contains the recombinant DNA vector of Claim 2 under 15 appropriate conditions such that expression of the
 - sequence is regulatable.

 5. A method of preparing a transformed host microorganism comprising a recombinant DNA molecule comprising a Streptomyces gal operon P2 promoter
- 20 expression unit or any functional derivative thereof and a foreign functional DNA sequence operatively linked to such expression unit, which comprises transforming an appropriate host microorganism or cell with such molecule.
- 6. A method of preparing a transformed host microorganism or cell comprising a recombinant DNA vector comprising the molecule of Claim 5 and, optionally, additionally comprising a replicon which comprises transforming an appropriate host microorganism with such vector.
- 7. A method of expressing a foreign functional DNA sequence which comprises cultivating a transformed microorganism or cell comprising a recombinant DNA vector comprising the molecule of Claim 5 and, optionally, additionally comprising a replicon, under suitable con-

ditions such that the functional DNA sequence is expressed.

- 8. A method of preparing a transformed host microorganism or cell comprising a recombinant DNA 5 molecule comprising a Streptomyces gal operon Pl promoter regulated region or any regulatable and functional derivative thereof and a foreign functional DNA sequence operatively linked to such regulated region. which comprises transforming an appropriate host micro-
- 9. A method of preparing a transformed host microorganism or cell comprising a recombinant DNA vector comprising the molecule of Claim 8 and, optionally, additionally comprising a replicon which comprises transforming an appropriate host microorganism or cell with such vector.
- DNA sequence which comprises cultivating a transformed host microorganism or cell comprising the recombinant 20 DNA vector of Claim 8 and, optionally, additionally
- comprising a replicon under suitable conditions such that the functional DNA sequence is expressed.
- 11. A method of regulating the expression of a foreign functional DNA sequence which comprises culti25 vating a transformed host microorganism or cell which contains a recombinant DNA vector comprising the molecule of Claim 8 and, optionally, additionally comprising a replicon, under appropriate conditions such that expression of the sequence is regulatable.
- 12. A method of preparing a transformed host microorganism or cell comprising a recombinant DNA molecule comprising a <u>Streptomyces gal</u> operon P2 promoter or any functional derivative thereof and a foreign functional DNA sequence operatively linked to the P2

promoter. which comprises transforming an appropriate host microorganism or cell with such molecule.

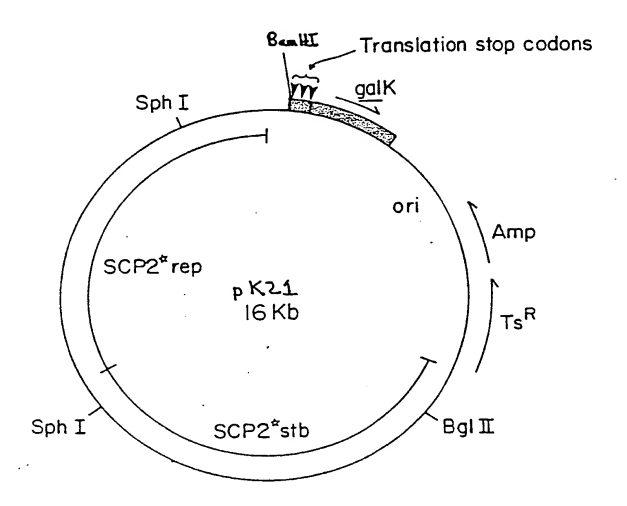
- microorganism or cell comprising a recombinant DNA vector comprising the molecule of Claim 12 and, optionally, additionally comprising a replicon, which comprises transforming an appropriate host microorganism
 with such vector.
- 14. A method of expressing a foreign functional

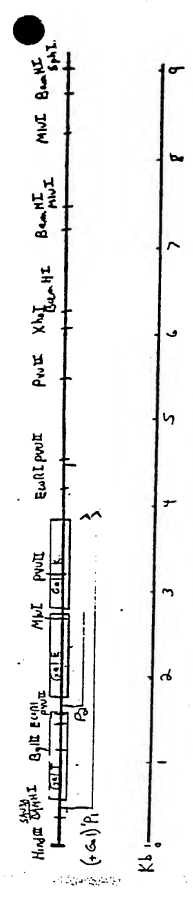
 10 DNA sequence which comprises cultivating a transformed host microorganism or cell comprising a recombinant DNA molecule of Claim 12 and, optionally, additionally comprising a replicon, under suitable conditions such that the functional DNA sequence is expressed.
- 15. A method of preparing a transformed host microorganism or cell comprising a recombinant DNA molecule comprising a <u>Streptomyces gal</u> operon Pl promoter or any regulatable and functional DNA sequence operatively linked to the Pl promoter, which comprises
- 20 transforming an appropriate host microorganism or cell with such molecule.
 - 16. A method of preparing a transformed host microorganism or cell comprising a recombinant DNA vector comprising the molecule of Claim 15 and, optio-
- 25 nally, additionally comprising a replicon, which comprises transforming an appropriate host microorganism with such vector.
 - 17. A method of expressing a foreign functional DNA sequence which comprises cultivating a transformed
- 30 host microorganism or cell comprising a recombinant DNA vector comprising the molecule of Claim 15 and, optionally, additionally comprising a replicon, under suitable conditions such that the functional DNA sequence is expressed.

- 18. A method of regulating the expression of a foreign functional DNA sequence which comprises cultivating a transformed host microorganism or cell which contains a recombinant DNA vector comprising the molecule of Claim 15 and, optionally, additionally comprising a replicon, under appropriate conditions such that expression of the sequence is regulatable.
- 19. A method of preparing a transformed host microorganism or cell comprising a recombinant DNA

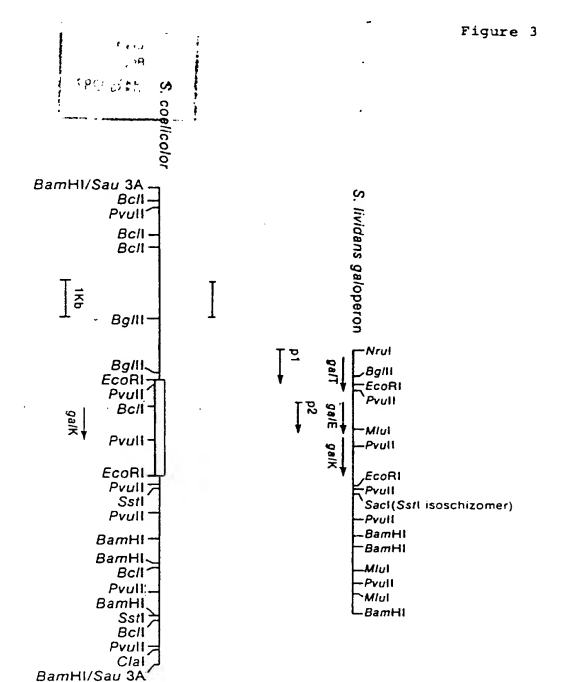
 10 molecule comprising a <u>Streptomyces gal</u> operon <u>galE</u> gene or any functional derivative thereof and a foreign functional DNA sequence operatively linked to the <u>galE</u> gene, which comprises transforming an appropriate host microorganism or cell with such molecule.
- microorganism or cell comprising a recombinant DNA molecule comprising a <u>Streptomyces gal</u> operon <u>gal</u>T gene or any functional derivative thereof and a foreign functional DNA sequence operatively linked to the <u>gal</u>T gene. which comprises transforming an appropriate host microorganism or cell with such molecule.
- 21. A method of preparing a transformed host microorganism or cell comprising a recombinant DNA molecule comprising a <u>Streptomyces lividans gal</u> operon 25 galk gene or any functional derivative thereof and a foreign functional DNA sequence operatively linked to the galk gene, which comprises transforming an appropriate host microorganism or cell with such molecule.
- 22. A method of enabling a nongalactose utilizing
 30 host microorganism or cell to utilize galactose which
 comprises transforming such host with a recombinant DNA
 vector or molecule comprising Streptomyces gal operon,
 or any portion of the Streptomyces gal operon which is
 adequate to enable such transformed host to utilize
 35 galactose, or any functional derivative thereof.

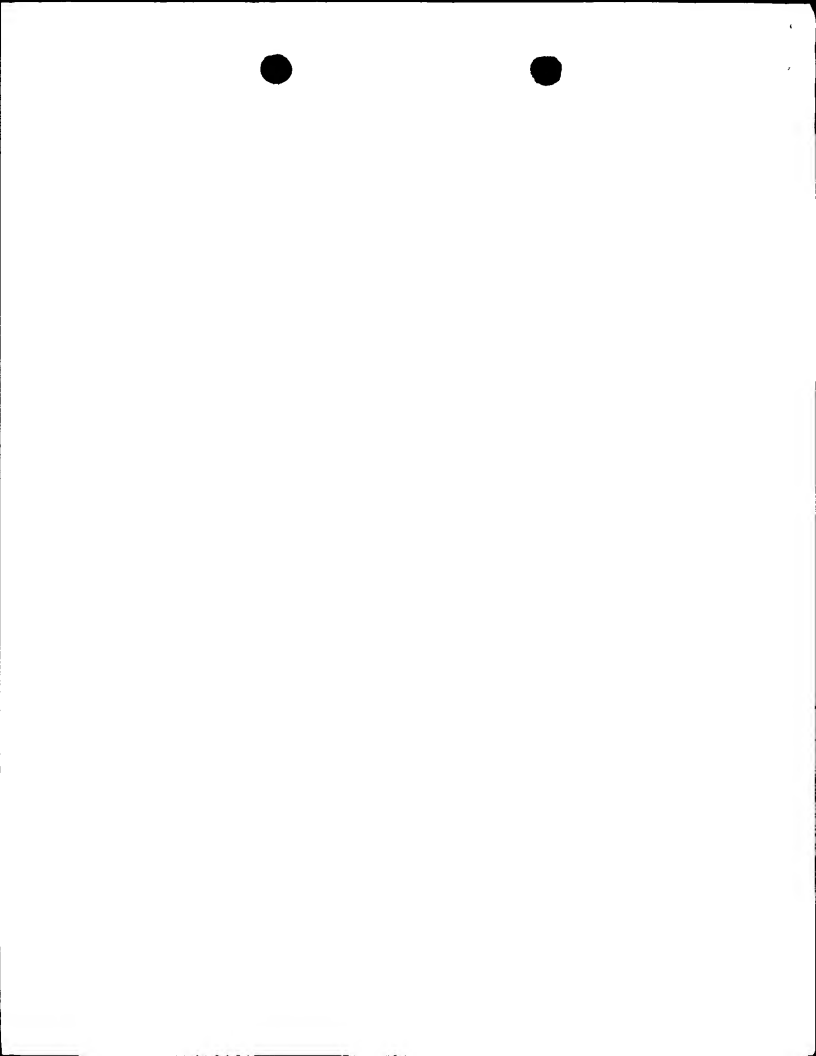
Figure 1





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0 235 112 A3

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(54) The gal operon of streptomyces.

(5) A recombinant DNA molecule comprising the *Streptomyces gal* operon *gal*K gene; *gal*E gene; *gal*T gene; P1 promoter; P2 promoter; P2 promoter expression unit; P1 promoter regulated region, or the entire *Streptomyces gal* operon is prepared.



EUROPEAN SEARCH REPORT

Application Number

EP 87 87 0026

	DOCUMENTS CONSIDE	RED TO BE RELEVANT					
Category	Citation of document with indica of relevant passage	tion, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl. 4)			
D,A	GENE, vol. 40, no. 2/3 191-201, Elsevier Scie Amsterdam, NL; M.E. BR "Characterization of S promoter sequences usi Escherichia coli galac	nce Publishers, AWNER et al.: treptomyces ng the		C 12 N 15/00 C 12 N 1/20			
D,A	NUCLEIC ACIDS RESEARCH 1985, pages 1841-1853, Oxford, GB; C. DEBOUCK "Structure of the gala Escherichia coli, the the gal operon"	IRL Press Ltd, (et al.: actokinase gene of					
Α	EP-A-0 187 630 (SMITH CORP.)	HKLINE BECKMAN					
A	ABSTRACTS OF THE ANNUA AMERICAN SOCIETY FOR MEAN, no. 0, 1984, abstract abstract abstract ansletional regulate the streptomyces - live operon"	MICROBIOLOGY, vol. ract no. H98; W. scriptional and ory elements in		TECHNICAL FIELDS SEARCHED (Int. Cl.4) C 12 N			
	The present search report has been	Date of completion of the search		Examiner			
TH	HE HAGUE	02-08-1988	PUL	AZZINI A.F.R.			
Y:p d A:t	CATEGORY OF CITED DOCUMENTS articularly relevant if taken alone articularly relevant if combined with anothe occument of the same category echnological background ion-written disclosure intermediate document	E : earlier patent doc after the filing da D : document cited fo L : document cited fo	T: theory or principle underlying th E: earlier patent document, but put after the filing date D: document cited in the application L: document cited for other reasons &: member of the same patent fam document				